By preliminary amendment, Applicants correct certain erroneous references to the sequence identifiers and add sequence identifiers where appropriate throughout the specification. These corrections and additions are shown in detail in the marked-up version of the affected paragraphs attached hereto as "Appendix - 1. "Revisions Made to Specification and Claims." Claims 7-21 and 34-37 have been canceled, and claims 3-6 and 24, 29, 30 and 32 have been amended. The changes to the amended claims are also shown in detail in Appendix - 1. A complete set of the pending claims, as presently amended, is attached as "Appendix - 2. Complete Set of Pending Claims, as Amended on September 17, 2001."

In the "Notice to File Missing Parts of Nonprovisional Application," the Initial Patent Examiner stated that the application is informal for the reason that excessive text is contained in the drawings as originally filed. In reply, substitute sheets of the drawings are submitted containing Figures 1-12 to replace the set of drawings filed on April 10, 2001. As required by the Initial Patent Examiner, the Figures have been revised to omit certain text so as to better comply with 37 CFR § 1.84. Text deleted from the Figures has been moved to the corresponding figure descriptions in the "Brief Description of the Drawings" section of the Specification, or has been moved to the "Detailed Description" section where the particular Figure is discussed, or has been deleted because it is considered duplicative of description in the Specification as originally filed. A marked-up copy of the Figures is attached as "Appendix-3. Details of Drawing Revisions Submitted on September 17, 2001," showing the proposed changes by circling the new text in red ink and by striking out deleted text in red ink. If the consent of the Examiner is necessary for entry of these substitute sheets of drawings, Applicants respectfully request approval by the Examiner of the proposed corrections to the Figures and entry of the substitute sheets of drawings.

Also enclosed is a computer diskette containing the initial computer readable form (CRF) of the Sequence Listing and a printed copy of the same, along with a statement that the content of the CRF and the original printed copy attached to the Application as filed is the same, and that the CRF and the attached printed copy are the same, and that no new matter is introduced thereby.

Applicants respectfully submit that the amendments to the specification, claims and drawings presented herein are to correct typographical errors, to facilitate examination of the application by ensuring compliance with 37 CFR § 1.821-1.824 or 37 CFR § 1.75(c) or 37 CFR § 1.84 prior to the first examination on the merits, or to comply with specific requirements stated in the "Notice to File Missing Parts." No new matter is introduced by way of these amendments.

If any additional fees are required or if the fees submitted are in excess of that required, please charge or credit the appropriate fees to Conley, Rose & Tayon, P.C. Deposit Account No.

03-2769/1889/00401/CGM. In the event that an extension of time is necessary in order for this submission to be deemed timely filed, please consider this a petition therefor and charge the associated fee to Conley, Rose & Tayon, P.C. Deposit Account No. 03-2769/1889/00401/CGM.

Respectfully submitted,

Carol G. Mintz

PTO Reg. No. 38,561

CONLEY, ROSE & TAYON

P.O. Box 3267

Houston, TX 77253-3267

(713) 238-8000 (Phone)

(713) 238-8008 (Fax)



APPENDIX - 1

REVISIONS MADE TO REPLACEMENT PARAGRAPHS AND CLAIMS

[0005] The invention relates to a compound that competitively inhibits binding of CSP [SEQ ID NO:4] to S. mutans histidine kinase [SEQ ID NO:6]. The compound is preferably a peptide or an antibody. The compound is preferably a derivative of [SEQ ID NO:2] [SEQ ID NO:3], a fragment of [SEQ ID NO:2] [SEQ ID NO:3] or a derivative of a fragment of [SEQ ID NO:2] [SEQ ID NO:3].

[0006] Schematic layout of the arrangement of the genetic locus encoding the signal peptide precursor (ComC) [SEQ ID NO:1], the histidine kinase (ComD) [SEQ ID NO:5] and the response regulator (ComE) [SEQ ID NO:7]. Note that this arrangement is different from other loci in related streptococci for the following reasons: a) The comC gene [SEQ ID NO:1] is transcribed from its own unique promoter, unlike the genes thus far described in other streptococci that are arranged in an operon-like cluster with the comC/DE genes being transcribed from a single promoter.

[0007] b) The comC gene [SEQ ID NO:1] is separated by 148 nucleotides from the comD gene [SEQ ID NO:5].

[0008] Shows the nucleic acid molecule that is [SEQ ID NO 1] [SEQ ID NO:3]. In a preferred embodiment, the figure shows CSP (competence signal peptide [SEQ ID NO 3]). Nucleotide sequence of the locus. Figure 2 also shows histidine kinase [SEQ ID NO 5] sequences and response regulator [SEQ ID NO 7] sequences.

[0008.5] Figure 2A. S. mutans comC gene [SEQ ID NO:1]. Encodes a precursor to a signal peptide [SEQ ID NO:2].

[0008.6] Figure 2B. S. mutans CSP encoding sequence [SEQ ID NO:3]. Encodes a Competence Signal Peptide [SEQ ID NO:4].

[0008.7] Figure 2C. S. mutans comD gene [SEQ ID NO:5]. Encodes a protein that functions as a histidine kinase receptor [SEQ ID NO:6].

[0008.8] Figure 2D. S. mutans comE gene [SEQ ID NO:7]. Encodes a response regulator that activates transcription of a number of genes [SEQ ID NO:8].

[[0009] Sequence of the deduced amino acid sequence of the signal peptide [SEQ ID NO 4], histidine kinase [SEQ ID NO: 6], and response regulator [SEQ ID NO: 8].

[0009.4] Figure 3A. S. mutans ComC protein (CSP Precursor) [SEQ ID NO:2].

[0009.5] Figure 3B. S. mutans ComD protein (Histidine Kinase) [SEQ ID NO:6].

[0009.6] Figure 3C. S. mutans ComE protein (Response Regulator) [SEQ ID NO:8].

- [0010] The deduced amino acid sequence of the signal peptide precursor <u>in various strains</u> and its predicted cleavage site, <u>following the glycine glycine signal</u> <u>The original peptide is expressed as a 46 amino acid peptide that is cleaved after the glycine-glycine residues to generate an active signal peptide</u>.
- [0011] Shows the peptide that is [SEQ ID NO:2] [SEQ ID NO:4]. The synthetic signal peptide [SEQ ID NO:16] that is effective at inducing competence, biofilm formation and acid tolerance in Streptococcus mutans.
- [0013] Table illustrating the effect of synthetic peptide on genetic competence in *S. mutans* cells. <u>Induction of genetic transformation in *Streptococcus mutans by synthetic* competence stimulating peptide (SCSP).</u>
- [0015] ComCDE local region [SEQ ID NO:23]. The ComC (first highlighted region; nucleotide 101 to 241), ComD (second highlighted region; nucleotides 383 to 1708) and ComE (third highlighted region; nucleotides 1705 to 2457) proteins are highlighted.
- [0016] The comX DNA sequence [SEQ ID NO:22], protein sequence [SEQ ID NO:23], and the comX gene local region [SEQ ID NO:24] with 100bp included both upstream and downstream (promoter is upstream).
- [0016.4] Figure 10A. S. mutans comX gene [SEQ ID NO:22].
- [0016.5] Figure 10B. S. mutans ComX protein [SEQ ID NO:25].
- [0016.6] Figure 10C. S. mutans comX gene local region [SEQ ID NO:26].
- [0017] The comA and comB nucleotide [SEQ ID NO:25] and [SEQ ID NO:27] and amino acid sequences [SEQ ID NO:26] and [SEQ ID NO:28]. ComA and ComB are the components of the CSP exporter.
- [0017.5] Figure 11A. S. mutans comA gene [SEQ ID NO:27].
- [0017.6] Figure 11B. S. mutans ComA protein [SEQ ID NO:28].
- [0017.7] Figure 11C. S. mutans comB gene [SEQ ID NO:29].
- [0017.8] Figure 11D. S. mutans ComB protein [SEQ ID NO:30].
- [0018] Illustrates the effect of synthetic peptide on acid resistance tolerance in *S. mutans* comC deficient cells. Addition of synthetic signal peptide (CSP) [SEQ ID NO:16] into the culture of the comC mutant restored the ability of the mutant to survive a low pH challenge when compared to the parent strain NG8.

[0020] We have identified a genetic locus in *S. mutans* consisting of three genes that encode:

1) a peptide precursor [SEQ ID NO:2] that is processed during export into a secreted 21-amino acid peptide (CSP) [SEQ ID NO:4]; 2) a histidine kinase [SEQ ID NO:6] that acts as a cell surface receptor activated by the peptide; 3) a response regulator [SEQ ID NO:8] that activates a number of other genes involved in genetic competence, biofilm formation, and acid tolerance of *S. mutans*. These properties have been attributed to the bacterium's ability to cause dental caries. Inactivation of any of these three genes or impairment of interaction or activity of any of their encoded proteins will disrupt the bacterium's ability to take up foreign DNA, form biofilms, and tolerate acidic pH.

Streptococcus mutans is a resident of the biofilm environment of dental plaque, a [0021] matrix of bacteria and extracellular material that adheres to the tooth surface. appropriate environmental conditions populations of S. mutans and the pH of the surrounding plaque will drop. S. mutans, being among the most acid tolerant organisms residing in dental plaque, will increase it numbers in this acidic environment and eventually become a dominant member of the plaque community. This situation eventually leads to dissolution of the tooth enamel, resulting in the development of dental caries. We control the accumulation and acid tolerance of this bacterium to make it less able to cause caries. We accomplish this by using inhibitors of an extracellular signal peptide that promotes the expression of genes involved in S. mutans biofilm formation and acid tolerance. The invention includes compounds that inhibit the action of the peptide. These inhibitors can include peptides, antibodies, or other agents that specifically inhibit the activation of the histidine kinase and the family of genes activated as a result of the histidine kinase activation by the signal molecule. Inhibitors include: modified structures of the peptide where amino acids are removed from the N- and/or COOH terminal of the peptide and/or substitutions of internal amino acid residues. We delete, one, two to 5, 6 to 10 and 10 to 15 amino acids from the peptide (for example at either terminal) and measure competitive inhibition of signal peptide binding to histidine

[0022] kinase (1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 or more amino acids are deleted and inhibition measured). Inhibitors also include antibodies raised against the 21-amino acid CSP [SEQ ID NO:4] alone or coupled to a larger molecule to increase immunogenicity.

[0024] In addition to identifying the genes encoding this signaling/sensing system, we have identified and chemically synthesized a 21-amino acid peptide [SEQ ID NO:16] that promotes biofilm formation and acid tolerance of *S. mutans*. A survey of the literature and genome databases reveals that genes similar to this signal-receptor system are present in most Gram-

positive bacteria, and therefore an inhibitor, or family of related inhibitors may be effective at inhibiting biofilm formation among a large group of bacteria.

[0025] The invention treats or prevents dental caries by addition of compounds that inhibit the stimulatory action of the 21-amino acid peptide [SEQ ID NO:4] on biofilm formation and acid tolerance of *S. mutans*. This is accomplished by delivery of these compounds to the biofilm and/or to incorporate these inhibitors into materials to control growth on surfaces. This includes delivery by topical application, alone or in combination with other compounds including toothpaste, mouthwash, food or food additives.

[0027] The invention includes an isolated CSP from *S. mutans*. The invention also includes a recombinant isolated CSP [SEQ ID NO:4] peptide produced by a cell including a nucleic acid molecule encoding CSP [SEQ ID NO:3] operably linked to a promoter. The invention also includes an isolated nucleic acid molecule encoding a CSP [SEQ ID NO:3]. The peptide we work with is preferably chemically synthesized [SEQ ID NO:16].

[0028] The invention includes CSP-encoding nucleic acid molecules [SEQ ID NO:3] and molecules having sequence identity or which hybridize to the CSP-encoding sequence and which encode a peptide having CSP activity (preferred percentages for sequence identity are described below) as well as vectors including these molecules. The invention also includes CSP [SEQ ID NO:4] or peptides having sequence identity (preferred percentages described below) or which have CSP activity. The nucleic acid molecules and peptides of the invention may be from S. mutans and they may be isolated from a native source, synthetic or recombinant. The invention includes CSP [SEQ ID NO:4] or peptides having sequence identity, which have CSP activity, as prepared by the processes described in this application.

[0029] The invention includes an isolated HK [SEQ ID NO:6] from *S. mutans*. The invention also includes a recombinant isolated HK polypeptide produced by a cell including a nucleic acid molecule encoding HK [SEQ ID NO:5] operably linked to a promoter. The invention also includes an isolated nucleic acid molecule encoding a HK polypeptide [SEQ ID NO:6].

[0030] The invention includes HK-encoding nucleic acid molecules and molecules having sequence identity or which hybridize to the HK-encoding sequence [SEQ ID NO:5] and which encode a protein having HK activity (preferred percentages for sequence identity are described below) as well as vectors including these molecules. The invention also includes HK [SEQ ID NO:4] or polypeptides having sequence identity (preferred percentages described below) or which have HK activity. The nucleic acid molecules and polypeptides of the invention may be

from *S. mutans* and they may be isolated from a native source, synthetic or recombinant. The invention includes HK [SEQ ID NO:4] or polypeptides having sequence identity, which have HK activity, as prepared by the processes described in this application.

[0031] The invention includes an isolated RR [SEQ ID NO:6] from S. mutans. The invention also includes a recombinant isolated RR [SEQ ID NO:6] polypeptide produced by a cell including a nucleic acid molecule encoding RR [SEQ ID NO:6] operably linked to a promoter. The invention also includes an isolated nucleic acid molecule encoding a RR polypeptide.

[0032] The invention includes RR-encoding nucleic acid molecules and molecules having sequence identity or which hybridize to the RR-encoding sequence [SEQ ID NO:5] and which encode a polypeptide having RR activity (preferred percentages for sequence identity are described below) as well as vectors including these molecules. The invention also includes RR [SEQ ID NO:6] or polypeptides having sequence identity (preferred percentages described below) or which have RR activity. The nucleic acid molecules and polypeptides of the invention may be from *S. mutans* and they may be isolated from a native source, synthetic or recombinant. The invention includes RR [SEQ ID NO:6] or polypeptides having sequence identity, which have RR activity, as prepared by the processes described in this application.

[0033] The comA and comB nucleotide [SEQ ID NO:27 and SEQ ID NO:29] and amino acid sequences [SEQ ID NO:28 and SEQ ID NO:30] are also aspects of the invention. ComA and ComB are components of the CSP exporter. The discussion of variants, sequence identity etc. for CSP, HK, RR applies to both the full sequences shown in the figures as well as bracketed portions of sequences (coding regions). The peptides and polypeptides may be natural, recombinantly produced or synthetic.

The invention includes nucleic acid molecules that are functional equivalents of all or part of the CSP sequence in [SEQ ID NO:1] [SEQ ID NO:3]. (A nucleic acid molecule may also be referred to as a DNA sequence or nucleotide sequence in this application. All these terms have the same meaning as nucleic acid molecule). Functionally equivalent nucleic acid molecules are DNA and RNA (such as genomic DNA, complementary DNA, synthetic DNA, and messenger RNA molecules) that encode peptides having the same or similar CSP activity as the CSP peptide shown in [SEQ ID NO:2] [SEQ ID NO:4]. Functionally equivalent nucleic acid molecules can encode peptides that contain a region having sequence identity to a region of a CSP peptide [SEQ ID NO:4] or more preferably to the entire CSP peptide. Identity is calculated according to methods known in the art. The ClustalW program (preferably using default parameters) [Thompson, JD et al., Nucleic Acid Res. 22:4673-4680.], described below, is most preferred. For example, if a nucleic acid molecule (called "Sequence A") has 90% identity to a

portion of the nucleic acid molecule in [SEQ ID NO:1] [SEQ ID NO:3], then Sequence A will preferably be identical to the referenced portion of the nucleic acid molecule in [SEQ ID NO:1] [SEQ ID NO:3], except that Sequence A may include up to 10 point mutations, such as substitutions with other nucleotides, per each 100 nucleotides of the referenced portion of the nucleic acid molecule in [SEQ ID NO:1] [SEQ ID NO:3]. Mutations described in this application preferably do not disrupt the reading frame of the coding sequence. Nucleic acid molecules functionally equivalent to the CSP sequences can occur in a variety of forms as described below.

[0035] Nucleic acid molecules may encode conservative amino acid changes in CSP peptide [SEQ ID NO:4]. The invention includes functionally equivalent nucleic acid molecules that encode conservative amino acid changes within a CSP amino acid sequence and produce silent amino acid changes in CSP.

[0036] Nucleic acid molecules may encode non-conservative amino acid substitutions, additions or deletions in CSP peptide. The invention includes functionally equivalent nucleic acid molecules that make non-conservative amino acid changes within the CSP amino acid sequence in [SEQ ID NO:2] [SEQ ID NO:4]. Functionally equivalent nucleic acid molecules include DNA and RNA that encode peptides, peptides and proteins having non-conservative amino acid substitutions (preferably substitution of a chemically similar amino acid), additions, or deletions but which also retain the same or similar CSP activity as the CSP peptide shown in [SEQ ID NO:2] [SEQ ID NO:4]. The DNA or RNA can encode fragments or variants of CSP. Fragments are useful as immunogens and in immunogenic compositions (U.S. Patent No. 5,837,472). The CSP or CSP-like activity of such fragments and variants is identified by assays as described below. Fragments and variants of CSP encompassed by the present invention should preferably have at least about 40%, 60%, 80% or 95% sequence identity to the naturally occurring CSP nucleic acid molecule, or a region of the sequence, such as the coding sequence or one of the conserved domains of the nucleic acid molecule, without being identical to the sequence in [SEQ ID NO:1] [SEQ ID NO:3]. Sequence identity is preferably measured with the ClustalW program (preferably using default parameters) (Thompson, JD et al., Nucleic Acid Res. 22:4673-4680)

[0037] Nucleic acid molecules functionally equivalent to the CSP nucleic acid molecule in [SEQ ID NO:1] [SEQ ID NO:3] will be apparent from the following description. For example, the sequence shown in [SEQ ID NO:1] [SEQ ID NO:3] may have its length altered by natural or artificial mutations such as partial nucleotide insertion or deletion, so that when the entire length of the coding sequence within [SEQ ID NO:1] [SEQ ID NO:3], is taken as 100%, the functional

equivalent nucleic acid molecule preferably has a length of about 60-120% thereof, more preferably about 80-110% thereof. Fragments may be less than 60%.

[0038] Nucleic acid molecules containing partial (usually 80% or less, preferably 60% or less, more preferably 40% or less of the entire length) natural or artificial mutations so that some codons in these sequences code for different amino acids, but wherein the resulting peptide retains the same or similar CSP activity as that of a naturally occurring CSP peptide [SEQ ID NO:4]. The mutated DNAs created in this manner should preferably encode a peptide having at least about 40%, preferably at least about 60%, at least about 80%, and more preferably at least about 90% or 95% sequence identity to the amino acid sequence of the CSP peptide in [SEQ ID NO:2] [SEQ ID NO:4]. The ClustalW program preferably assesses sequence identity.

[0039] Since the genetic code is degenerate, the nucleic acid sequence in [SEQ ID NO:1] [SEQ ID NO:3] is not the only sequence which may code for a peptide having CSP activity. This invention includes nucleic acid molecules that have the same essential genetic information as the nucleic acid molecule described in [SEQ ID NO:1] [SEQ ID NO:3]. Nucleic acid molecules (including RNA) having one or more nucleic acid changes compared to the sequences described in this application and which result in production of a peptide shown in [SEQ ID NO:2] [SEQ ID NO:4] are within the scope of the invention.

[0040] Other functional equivalent forms of CSP-encoding nucleic acids can be isolated using conventional DNA-DNA or DNA-RNA hybridization techniques. Thus, the present invention also includes nucleic acid molecules that hybridize to one or more of the sequences in [SEQ ID NO:1] [SEQ ID NO:3] or its complementary sequence, and that encode expression for peptides, peptides and proteins exhibiting the same or similar activity as that of the CSP peptide produced by the DNA in [SEQ ID NO:1] [SEQ ID NO:3] or its variants. Such nucleic acid molecules preferably hybridize to the sequence in [SEQ ID NO:1] [SEQ ID NO:3] under moderate to high stringency conditions (see Sambrook et al. Molecular Cloning: A Laboratory Manual, Most Recent Edition, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.). High stringency washes have low salt (preferably about 0.2% SSC), and low stringency washes have high salt (preferably about 2% SSC). A temperature of about 37 °C or about 42 °C is considered low stringency, and a temperature of about 50-65 °C is high stringency. The invention also includes a method of identifying nucleic acid molecules encoding a CSP activator peptide (preferably a mammalian peptide), including contacting a sample containing nucleic acid molecules including all or part of [SEQ ID NO:1] [SEQ ID NO:3] (preferably at least about 15 or 20 nucleotides of [SEQ ID NO:1] [SEQ ID NO:3]) under moderate or high stringency hybridization conditions and identifying nucleic acid molecules which hybridize to the nucleic

acid molecules including all or part of [SEQ ID NO:1] [SEQ ID NO:3].). Similar methods are described in U.S. Patent No. 5,851,788, which is incorporated by reference in its entirety.

[0041] The invention also includes methods of using all or part of the nucleic acid molecules which hybridize to all or part of [SEQ ID NO:1] [SEQ ID NO:3], for example as probes or in assays to identify antagonists or inhibitors of the peptides produced by the nucleic acid molecules (described below). The invention also includes methods of using nucleic acid molecules having sequence identity to the CSP nucleic acid molecule [SEQ ID NO:3] (as described below) in similar methods.

[0042] The invention also includes a nucleic acid molecule detection kit including, preferably in a suitable container means or attached to a surface, a nucleic acid molecule of the invention encoding CSP [SEQ ID NO:4] or a peptide having CSP activity and a detection reagent (such as a detectable label). Other variants of kits will be apparent from this description and teachings in patents such as U.S. Patent Nos. 5,837,472 and 5,801,233,which are incorporated by reference in their entirety.

A nucleic acid molecule described above is considered to have a function substantially equivalent to the CSP nucleic acid molecules [SEQ ID NO:3] of the present invention if the peptide [SEQ ID NO:4] produced by the nucleic acid molecule has CSP activity. A peptide has CSP activity if it can stimulate genetic competence and acid tolerance in S. mutans. Activation of the HK [SEQ ID NO:6]/RR [SEQ ID NO:8] is shown where a peptide is capable of stimulating the uptake and incorporation of foreign DNA. We describe below how the activity of these peptide-mediated processes can be measured by determining the efficiency of plasmid uptake, which is a measure of genetic competence. Since the ability to transport and incorporate foreign DNA relies on activation of the HK [SEQ ID NO:6]/RR [SEQ ID NO:8] and subsequent genes activated by the signal cascade initiated by the signal peptide, measurement of the conferment of erythromycin resistance by cells exposed to the peptide and plasmid DNA conferring erythromycin resistance indicates its level of function. Conversely if an inhibitor is capable of interfering with the action of the peptide the competence assay will indicate this by a corresponding decrease in the number of cells that acquire erythromycin resistance as described in the assays below (assays of genetic competence and assay of transformation of biofilms). Activation of the HK [SEQ ID NO:6]/RR [SEQ ID NO:8] is also shown where a peptide is capable of stimulating an acid tolerance response. We describe below how the activity of these peptide-mediated processes can be measured by determining the survival rate of cells in acidic pH conditions. Since the ability to survive exposure to acidic pH depends on the activation of the HK/RR and subsequent genes activated by the signal peptide, measurement of the survival of S. mutans in low pH conditions indicates the level of function of

the signal peptide. Conversely, if an inhibitor is capable of interfering with the signal peptide sensing system the assay for acid adaptation will indicate this by a corresponding decrease in the survival rate of cells grown in acidic pH conditions as described in the assay below (assay of acid adaptation).

[0049] Another embodiment of the invention relates to a method of transfecting a cell with a nucleic acid molecule of the invention, inserted in an expression vector to produce a cell expressing the CSP peptide [SEQ ID NO:4] or other peptide of the invention. The invention also relates to a method of expressing the peptides of the invention in a cell. A preferred process would include culturing a cell including a recombinant DNA vector including a nucleic acid molecule encoding CSP [SEQ ID NO:3] (or another nucleic acid molecule of the invention) in a culture medium so that the peptide is expressed. The process preferably further includes recovering the peptide from the cells or culture medium.

[0050] The invention also includes oligonucleotide probes made from the cloned CSP nucleic acid molecules described in this application or other nucleic acid molecules of the invention (see Materials and Methods section). The probes may be 15 to 20 nucleotides in length. A preferred probe is at least 15 nucleotides of CSP in [SEQ ID NO:1] [SEQ ID NO:3]. The invention also includes at least 15 consecutive nucleotides of [SEQ ID NO:1] [SEQ ID NO:3]. The probes are useful to identify nucleic acids encoding CSP peptides as well as peptides functionally equivalent to CSP. The oligonucleotide probes are capable of hybridizing to the sequence shown in [SEQ ID NO:1] [SEQ ID NO:3] under stringent hybridization conditions. A nucleic acid molecule encoding a peptide of the invention may be isolated from other organisms by screening a library under moderate to high stringency hybridization conditions with a labeled probe. The activity of the peptide encoded by the nucleic acid molecule is assessed by cloning and expression of the DNA. After the expression product is isolated, the peptide is assayed for CSP activity as described in this application.

[0051] Functionally equivalent CSP nucleic acid molecules from other cells, or equivalent CSP-encoding cDNAs or synthetic DNAs, can also be isolated by amplification using Polymerase Chain Reaction (PCR) methods. Oligonucleotide primers, such as degenerate primers, based on [SEQ ID NO:1] [SEQ ID NO:3] can be prepared and used with PCR and reverse transcriptase (E. S. Kawasaki (1990), In Innis et al., Eds., PCR Protocols, Academic Press, San Diego, Chapter 3, p. 21) to amplify functional equivalent DNAs from genomic or cDNA libraries of other organisms. The oligonucleotides can also be used as probes to screen cDNA libraries.

The present invention includes not only the peptides encoded by the sequences of the [0052] invention, but also functionally equivalent peptides, peptides and proteins that exhibit the same or similar CSP peptide activity. A peptide is considered to possess a function substantially equivalent to that of the CSP peptide [SEQ ID NO:4] if it has CSP activity. CSP activity means that it is able to confer genetic competence to S. mutans, as measured by an increased ability to incorporate and express foreign genetic material, when added to cells as described in the assay of genetic competence below. CSP activity also means that the peptide is able to confer an acid tolerance response in S. mutans as measured by an increase in cell survival under acidic pH conditions when added to cells as described in the assay for acid adaptation below. Functionally equivalent peptides, peptides and proteins include peptides, peptides and proteins that have the same or similar protein activity as CSP when assayed, i.e. they are able to stimulate genetic competence and low pH tolerance (the ability to withstand acid challenges of pH 3.5 -pH 3.0 for up to 3 hours) in S. mutans. A peptide has CSP activity if it is capable of increasing the frequency of uptake and expression of foreign DNA as described in the following assay for genetic competence and if the peptide can promote an acid tolerance response as described in the assay for acid adaptation.

[0053] Identity refers to the similarity of two peptides or proteins that are aligned so that the highest order match is obtained. Identity is calculated according to methods known in the art, such as the ClustalW program. For example, if a peptide (called "Sequence A") has 90% identity to a portion of the peptide in [SEQ ID NO:2] [SEQ ID NO:16], then Sequence A will be identical to the referenced portion of the peptide in [SEQ ID NO:2] [SEQ ID NO:16], except that Sequence A may include up to 1 point mutations, such as substitutions with other amino acids, per each 10 amino acids of the referenced portion of the peptide in [SEQ ID NO:2] [SEQ ID NO:2] [SEQ ID NO:16]. Peptides, peptides and proteins functional equivalent to the CSP peptides can occur in a variety of forms as described below.

[0054] Peptides biologically equivalent in function to CSP peptide include amino acid sequences containing amino acid changes in the CSP sequence [SEQ ID NO:4]. The functional equivalent peptides have at least about 40% sequence identity, preferably at least about 60%, at least about 75%, at least about 80%, at least about 90% or at least about 95% sequence identity, to the natural CSP peptide [SEQ ID NO:4] or a corresponding region. The ClustalW program preferably determines sequence identity. Most preferably, 1, 2, 3, 4, 5, 5-10, 10-15 amino acids are modified.

[0058] The variants preferably retain the same or similar CSP activity as the naturally occurring CSP [SEQ ID NO:4]. The CSP activity of such variants can be assayed by techniques described in this application and known in the art.

[0059] Variants produced by combinations of the techniques described above but which retain the same or similar CSP activity as naturally occurring CSP [SEQ ID NO:4] are also included in the invention (for example, combinations of amino acid additions, and substitutions).

[0062] As well, probes and antibodies for a histidine kinase [SEQ ID NO:5 and SEQ ID NO:6], response regulator [SEQ ID NO:7 and SEQ ID NO:8] comA [SEQ ID NO:27 and SEQ ID NO:28] or comB [SEQ ID NO:29 and SEQ ID NO:30] may be prepared using the description in this application and techniques known in the art. The description for preparation of CSP variants and mutants is also applicable to the histidine kinase [SEQ ID NO:5 and SEQ ID NO:6], response regulator [SEQ ID NO:7 and SEQ ID NO:8] or comA [SEQ ID NO:27 and SEQ ID NO:28] and comB [SEQ ID NO:29 and SEQ ID NO:30] of the invention. The invention also includes fragments of HK [SEQ ID NO:5 and SEQ ID NO:6] having HK activity, fragments of RR [SEQ ID NO:7 and SEQ ID NO:8] having RR activity and fragments of comA [SEQ ID NO:28] or comB [SEQ ID NO:29 and SEQ ID NO:30] having activity.

[0063] The activity of the CSP peptide [SEQ ID NO:4] may be varied by carrying out selective site-directed mutagenesis. We characterize the binding domain and other critical amino acid residues in the peptide that are candidates for mutation, insertion and/or deletion. Sequence variants may be synthesized. A DNA plasmid or expression vector containing the CSP nucleic acid molecule [SEQ ID NO:3] or a nucleic acid molecule having sequence identity may be used for these studies using the U.S.E. (Unique site elimination) mutagenesis kit from Pharmacia Biotech or other mutagenesis kits that are commercially available, or using PCR. Once the mutation is created and confirmed by DNA sequence analysis, the mutant peptide is expressed using an expression system and its activity is monitored. This approach is useful to identify CSP inhibitors. All these modifications of the CSP DNA sequences [SEQ ID NO:3] presented in this application and the peptides produced by the modified sequences are encompassed by the present invention.

[0064] The CSP inhibitors are also useful when combined with a carrier in a pharmaceutical composition. The compositions are useful when administered in methods of medical treatment or prophylaxis of a disease, disorder or abnormal physical state caused by *S. mutans*. The invention also includes methods of medical treatment of a disease, disorder or abnormal

physical state characterized by excessive *S. mutans* or levels or activity of CSP peptide **[SEQ ID NO:4]**, for example by administering a pharmaceutical composition including a carrier and a CSP inhibitor. Caries is one example of a disease, which can be treated or prevented by antagonizing CSP **[SEQ ID NO:4]**.

[0066] CSP activity could be blocked by antisense mRNA or by inhibiting the activity of the exporter that secretes it from the cell. We have the sequence of these exporters. There are two copies of the genes (comAB) [SEQ ID NO:27 and SEQ ID NO:29] that are involved in export.

[0067] Nucleic acid molecules (antisense inhibitors of CSP) [SEQ ID NO:3] and competitive inhibitors of CSP [SEQ ID NO:4] may be introduced into cells using *in vivo* delivery vehicles such as liposomes. They may also be introduced into these cells using physical techniques such as microinjection and electroporation or chemical methods such as coprecipitation or using liposomes.

[0070] Antibodies directed against the CSP [SEQ ID NO:4 or SEQ ID NO:16] would provide protection against caries. Antibodies may be manufactured as described below. Alternatively, a peptide of the invention [SEQ ID NO:4 or SEQ ID NO:16] or a fragment thereof may be used with a carrier to make a vaccine. The peptide or fragment may also be conjugated to another molecule to increase its antigenicity. Antibodies can also be coupled to the peptide (Brady, L.J. et al., "Monoclonal Antibody-Mediated Modulation of the Humoral Immune Response against Mucosally Applied Streptococcus mutans" (in press). In order to enhance the immune response the peptide can be coupled to KLH, ovalbumin, or thyroglobulin prior to immunization. The vaccine composition will trigger the mammal's immune system to produce antibodies. The invention includes vaccine compositions and methods of vaccinating a mammal, preferably a human, against dental caries by administering to the mammal an effective amount of a vaccine composition. Techniques for preparing and using vaccines are known in the art. To prepare the vaccine, the peptide, or a fragment of the peptide, may be mixed with other antigens (of different immunogenicity), a vehicle or an excipient. Examples of peptide vaccines are found in U.S. Patent Nos. 5,679,352, 5,194,254 and 4,950,480. Techniques for preparing vaccines involving site-directed mutagenesis are described in U.S. Patent Nos. 5,714,372, 5,543,302, 5,433,945, 5,358,868, 5,332,583, 5,244,657, 5,221,618, 5,147,643, 5,085,862 and 5,073,494. Vaccines may be administered by known techniques, such as topical or parenteral administration. Vast changes are taking place in vaccinology consequent to the introduction of new technologies. Acellular purified fractions devoid of side effects, non-pathogenic but immunogenic mutants, recombinant technology, conjugated vaccines, combination vaccines (to limit the number of injections). Vaccine delivery systems can deliver multiple doses of the vaccine at a single

contact point. A genetically engineered oral vaccine is useful to impart better and longer duration of immunity. Oral vaccines are useful. The nose as a route for immunization is also useful. DNA alone can constitute the vaccines, inducing both humoral and cell-mediated immune responses. Live recombinant vaccines are also useful. Potent adjuvants add to the efficacy of the vaccines. One can also 'humanize' mouse monoclonals by genetic engineering and express these efficiently in plants. These recombinant antibodies are opening out an era of highly specific and safe therapeutic interventions. An advantage of preformed antibodies directed at a defined target and given in adequate amounts is the certainty of efficacy in every recipient, in contrast to vaccines, where the quality and quantum of immune response varies from individual to individual. For example, nasal immunization may be done as described in C. Jespersgaard et al. "Protective Immunity against *Streptococcus mutans* Infection in Mice after Intranasal Immunization with the Glucan-Binding Region of *S. mutans* Glucosyltransferase" Infection and Immunity, December 1999, p. 6543-6549, Vol. 67, No. 12. Vaccine compositions may comprise solid or liquid formulations such as gels, sprays, inhalants, tablets, toothpastes, mouthwashes or chewing qum.

[0072] Inhibitors are preferably directed towards CSP [SEQ ID NO:4 or SEQ ID NO:16] to block *S. mutans* competence, low pH tolerance and biofilm formation.

[0073] A method of identifying a compound which reduces the interaction of CSP [SEQ ID NO:4 or SEQ ID NO:16] with HK [SEQ ID NO:6], can include: contacting (i) CSP [SEQ ID NO:4 or SEQ ID NO:16] with (ii) HK [SEQ ID NO:6], a CSP-binding fragment of HK [SEQ ID NO:6] or a derivative of either of the foregoing in the presence of the compound; and b)

determining whether the interaction between (i) and (ii) is reduced, thereby indicating that the compound reduces the interaction of CSP [SEQ ID NO:4 or SEQ ID NO:16] and HK [SEQ ID NO:6]. A CSP inhibitor (caries treating or preventing compound) inhibits the interaction between (i) and (ii). By way of example, one can screen a synthetic peptide library. One could also screen small non-peptide organic molecules.

[0074] In one embodiment, the invention includes an assay for evaluating whether test compounds are capable of acting as agonists or antagonists for CSP [SEQ ID NO:4], or a peptide having CSP functional activity, including culturing cells containing DNA which expresses CSP [SEQ ID NO:3], or a peptide having CSP activity so that the culturing is carried out in the presence of at least one compound whose ability to modulate CSP activity is sought to be determined and thereafter monitoring the cells for either an increase or decrease in the level of CSP [SEQ ID NO:4 or SEQ ID NO:16] or CSP activity. Other assays (as well as variations of

the above assay) will be apparent from the description of this invention and techniques such as those disclosed in U.S. Patent No. 5,851,788, 5,736,337 and 5,767,075 which are incorporated by reference in their entirety. For example, the test compound levels may be either fixed or variable.

[0075] The CSP [SEQ ID NO:4 or SEQ ID NO:16] peptide is also useful as an antigen for the preparation of antibodies that can be used to purify or detect other CSP-like peptides. Antibodies may also block CSP [SEQ ID NO:4] binding to HK [SEQ ID NO:6]. Antibodies are preferably targeted to the entire CSP [SEQ ID NO:4] sequence. The CSP peptide [SEQ ID NO:4 or SEQ ID NO:16] may be conjugated to other compounds, in order to increase immunogenicity.

[0076] We generate polyclonal antibodies against the CSP [SEQ ID NO:4 or SEQ ID NO:16], which is a unique sequence. Monoclonal and polyclonal antibodies are prepared according to the description in this application and techniques known in the art. For examples of methods of preparation and uses of monoclonal antibodies, see U.S. Patent Nos. 5,688,681, 5,688,657, 5,683,693, 5,667,781, 5,665,356, 5,591,628, 5,510,241, 5,503,987, 5,501,988, 5,500,345 and 5,496,705, which are incorporated by reference in their entirety. Examples of the preparation and uses of polyclonal antibodies are disclosed in U.S. Patent Nos. 5,512,282, 4,828,985, 5,225,331 and 5,124,147 which are incorporated by reference in their entirety. Antibodies recognizing CSP [SEQ ID NO:4 or SEQ ID NO:16] can be employed to screen organisms or tissues containing CSP peptide [SEQ ID NO:4] or CSP-like peptides. The antibodies are also valuable for immuno-purification of CSP [SEQ ID NO:4] or CSP-like peptides from crude extracts.

[0077] An antibody (preferably the antibody described above) may be used to detect CSP [SEQ ID NO:4] or a similar peptide, for example, by contacting a biological sample with the antibody under conditions allowing the formation of an immunological complex between the antibody and a peptide recognized by the antibody and detecting the presence or absence of the immunological complex whereby the presence of CSP [SEQ ID NO:4] or a similar peptide is detected in the sample. The invention also includes compositions preferably including the antibody, a medium suitable for the formation of an immunological complex between the antibody and a peptide recognized by the antibody and a reagent capable of detecting the immunological complex to ascertain the presence of CSP [SEQ ID NO:4] or a similar peptide. The invention also includes a kit for the *in vitro* detection of the presence or absence of CSP [SEQ ID NO:4] or a similar peptide in a biological sample, wherein the kit preferably includes an antibody, a medium suitable for the formation of an immunological complex between the antibody and a peptide recognized by the antibody and a reagent capable of detecting the

immunological complex to ascertain the presence of CSP [SEQ ID NO:4] or a similar peptide in a biological sample. Further background on the use of antibodies is provided, for example in U.S. Patent Nos. 5,695,931 and 5,837,472, which are incorporated by reference in their entirety. The ability of the peptide to activate the HK [SEQ ID NO:6] and RR [SEQ ID NO:8] and the subsequent genes involved in the conferral of the properties of genetic competence, acid tolerance and biofilm formation can be determined by measuring the efficiency of uptake and expression of DNA (preferably plasmid DNA) in S. mutans when exposed to signal peptide and/or inhibitor. Two methods modified based on the protocols described by Perry et al. Infect Immun, 41:722-727 and Lindler and Macrina J Bacteriol, 166:658-665 are used to assay genetic competence. The method involves adding DNA and CSP [SEQ ID NO:3] (preferably plasmid DNA) to a S. mutans culture (or culture of a bacteria expressing CSP [SEQ ID NO:3] or a variant thereof). The rate of transformation is then determined. S. mutans is preferably grown in THYE plus 5% horse serum (THYE-HS). After 2-hr incubation, 1 μg/ml plasmid DNA or 10 μg/ml of chromosomal DNA is added to the culture. To assay induction of competence, competence signal peptide, (SCSP) [SEQ ID NO:16] is then added to the cultures, incubation continued for 30 minutes with a final concentration of 500 ng/ml of SCSP added to each sample. After the 30minute incubation equal amounts of DNA is added to each well (1 µg/ml plasmid or 10 µg/ml of chromosomal DNA) and incubation continued for another 2 hrs.. Cell dilutions were immediately spread on THYE agar plates plus appropriate antibiotics. Transformation frequency was expressed as the number of transformants (antibiotic resistant cells) per number of viable recipients. This is determined by comparing the number of cells able to grow in the presence of antibiotic (conferred by the applied plasmid or chromosomal DNA) relative to the total number of cells present (i.e., that grow in the absence of antibiotic). A higher value indicates a higher rate of transformation and thus is reflective of a stimulatory effect by the peptide. Consequently, addition of a molecule that successfully acts as an inhibitor results in a lower ratio of transformants/recipients, indicating that the inhibitor is effective at blocking activity of the CSP [SEQ ID NO:4]. CSP deficient cells [SEQ ID NO:3 or SEQ ID NO:4] may also be used in a variation of these assays. One can identify compounds that inhibit CSP [SEQ ID NO:4] or variants thereof by adding a test compound to the mixture to determine if the rate of transformation is decreased by the addition of the test compound.

[0079] The activity of the system can also be measured by an *in vitro* assay that relies on the measurement of marker protein expression (such as green fluorescent protein (GFP)) via expression from a fusion to a promoter controlled by the signal cascade initiated by CSP [SEQ ID NO:4]/HK [SEQ ID NO:6]/RR [SEQ ID NO:8]. One such promoter occurs immediately 5' proximal to the *S. mutans* comX gene. *S. mutans* cells grown in microtiter wells are exposed to

the CSP [SEQ ID NO:4] and/or inhibitor and the level of fluorescence of the comX::GFP strain is measured to give a quantitative measure of CSP [SEQ ID NO:4] stimulation (and conversely inhibitor activity). One can identify compounds that inhibit CSP [SEQ ID NO:4] or variants thereof by adding a test compound to the mixture to determine if the quantitative measure of CSP [SEQ ID NO:4] stimulation is decreased by the addition of the test compound.

The ability of CSP [SEQ ID NO:4] to promote acid resistance tolerance is determined [0800] by measuring the cell survival rate of S. mutans when exposed to acidic pH. In one example, S. mutans are first grown in batch culture to assay acid tolerance response in 'standard' log- and stationary-phase cells by using a modification of methods described previously by Svensäter et al. Oral Microbiol. Immunol., 12:266-73. Mid-log-phase cells are obtained by transferring one volume of overnight culture into nine volumes (1:10) of fresh TYG medium (pH 7.5) and incubated at 37°C with 5% CO2 for 2 hours. These cells are then collected by centrifugation at 8,000 x g for 10 min and resuspended in 2 ml of fresh TYG (pH 5.5) at various cell densities as determined by O.D₆₀₀. The cells are induced for acid adaptation by incubation at pH 5.5 for 2 h at 37°C with 5% CO2. The adapted log-phase cells are then exposed to the killing pH. Killing pH is pre-determined by incubating unadapted, mid-log phase cells in TYG medium at pH values from 6.0 to 2.0. Stationary-phase cells are prepared by re-suspending late-log phase cells in TY medium (tryptone-yeast extract) without glucose. The culture is incubated at 37°C for 2 h to allow the cells to fully enter into stationary phase. Induction of acid adaptation in stationary-phase cells follows a similar procedure to that for log-phase cells. Adaptation of both log- and stationary-phase cells to acidic pH is determined by measuring the ability of bacterial cells to survive a killing pH for 3 h. Acid killing is initiated by resuspending cells in the same volume of fresh TYG (pH 3.5) and an aliquot of cell suspension is taken immediately from each sample to determine total viable cell number at zero time. The cells are then incubated for 3 h at 37°C with 5% CO2 and an aliquot of sample is taken to determine survival rate by viable cell counts. Addition of a molecule that successfully acts as an inhibitor results in a decrease in the acid resistance tolerance of S mutans resulting in a corresponding decrease in cell survival indicating that the inhibitor is effective at blocking activity of CSP [SEQ ID NO:4]. CSP [SEQ ID NO:3 or SEQ ID NO:4] deficient cells may also be used in a variation of these assays wherein addition of the signal peptide can complement the acid-adaptation-defective phenotype of a comC [SEQ ID NO:1 or SEQ ID NO:2] deficient cell. One can identify compounds that inhibit CSP [SEQ ID NO:3 or SEQ ID NO:4] or variants thereof by adding a test compound to the mixture to determine if the survival rate of cells is decreased by the addition of the test compound

Cells transformed with a nucleic acid molecule of the invention (histidine kinase [SEQ [1800] ID NO:51, CSP [SEQ ID NO:3] or response regulator [SEQ ID NO:7]) are useful as research tools. For example, one may obtain a cell (or a cell line, such as an immortalized cell culture or a primary cell culture) that does not express histidine kinase [SEQ ID NO:5], CSP [SEQ ID NO:3] or response regulator [SEQ ID NO:8], insert a histidine kinase [SEQ ID NO:5], CSP [SEQ ID NO:3] or response regulator [SEQ ID NO:7] nucleic acid molecule in the cell, and assess the level of expression and activity. Alternatively, histidine kinase [SEQ ID NO:5], CSP [SEQ ID NO:3] or response regulator [SEQ ID NO:7] nucleic acid molecules may be over-expressed in a cell that expresses a histidine kinase [SEQ ID NO:5], CSP [SEQ ID NO:3] or response regulator [SEQ ID NO:7] nucleic acid molecule. In another example, experimental groups of cells may be transformed with vectors containing different types of histidine kinase [SEQ ID NO:5], CSP [SEQ ID NO:3] or response regulator [SEQ ID NO:7] nucleic acid molecules to assess the levels of polypeptides and peptides produced, its functionality and the phenotype of the cells. The polypeptides and peptides are also useful for in vitro analysis of histidine [SEQ ID NO:6] kinase, CSP [SEQ ID NO:4] or response regulator [SEQ ID NO:8] activity or structure. For example, the polypeptides and peptides produced can be used for microscopy or X-ray crystallography studies.

The histidine kinase [SEQ ID NO:5 and SEQ ID NO:6], CSP [SEQ ID NO:3 and SEQ [0082] ID NO:4] or response regulator [SEQ ID NO:7 and SEQ ID NO:8] nucleic acid molecules and polypeptides are also useful in assays for the identification and development of compounds to inhibit and/or enhance polypeptide or peptide function directly. For example, they are useful in an assay for evaluating whether test compounds are capable of acting as antagonists for histidine kinase [SEQ ID NO:6], CSP [SEQ ID NO:4] or response regulator [SEQ ID NO:8] by: (a) culturing cells containing a nucleic acid molecule which expresses histidine kinase [SEQ ID NO:5], CSP [SEQ ID NO:3] or response regulator peptides [SEQ ID NO:7] (or fragments or variants thereof having histidine [SEQ ID NO:6] kinase, CSP or response regulator activity) wherein the culturing is carried out in the presence of increasing concentrations of at least one test compound whose ability to inhibit histidine [SEQ ID NO:6] kinase, CSP [SEQ ID NO:4] or response regulator [SEQ ID NO:8] is sought to be determined; and (b) monitoring in the cells the level of inhibition as a function of the concentration of the test compound, thereby indicating the ability of the test compound to inhibit histidine kinase [SEQ ID NO:6], CSP [SEQ ID NO:4] or response regulator [SEQ ID NO:8] activity.

[0085] Biofilms are developed on polystyrene microtiter plates to provide a rapid and simple method for assaying biofilm formation, and hence activity of the peptide [SEQ ID NO:4]/receptor

[SEQ ID NO:8]/kinase **[SEQ ID NO:6]** system. Formation of biofilms is initiated by inoculating 20 ul of cell suspension into each well containing 2 ml of biofilm medium (4X diluted Todd-Hewitt Yeast Extract supplemented with final concentration of 0.01% hog gastric mucin) for overnight incubation at 37°C under an anaerobic condition. After 20-h incubation, fluid medium is removed and added with 2 ml of pre-warmed, fresh THYE plus 5% horse serum. The cultures are incubated for 30 minutes and each well is supplemented with a final concentration of 200 ng/ml of synthetic competence stimulating peptide (SCSP) and varying concentrations of the inhibitor and the incubation is continued. After 30 minutes, plasmid DNA (1 mg/ml) or chromosomal DNA (10 mg/ml) is added to each well and the cultures are incubated for an additional 2 hr. Planktonic cells are then removed and the wells are washed once with PBS buffer. Biofilm cells are collected into 2 ml fresh medium by a gentle sonication or washing the wells using a pipette. The samples are centrifuged at 12,000 x g for 5 min. Both biofilm and planktonic cells are resuspended into 200 μ l of fresh medium and are immediately spread on THYE agar plus appropriate antibiotics. Transformation frequency is determined after 48-h of incubation.

[0086] Homologues of the Streptococcus pneumoniae comD [SEQ ID NO:5]/E [SEQ ID NO:5]/E [SEQ ID NO:6]/ response regulator [SEQ ID NO:8] system were identified. This sequence was used to design primers to amplify the region from a number of S. mutans isolates. An open reading frame consisting of 138 nucleotides was located 148 nucleotides 5' proximal from the end of the comD homolog in the opposite orientation (Fig 1). This ORF was found to encode a peptide of 46-amino acid [SEQ ID NO:2] in length, the precursor of the 21-amino acid CSP [SEQ ID NO:4].

[0087] The comCDE genes [SEQ ID NO:23] were amplified from the genomes of several S. mutans isolates by PCR using primers designed based on the genome database sequence and their nucleotide sequences determined. The deduced amino acid sequences are compared among the isolates by sequence alignment to confirm identity.

[0088] Genes are inactivated by integration of internal homologous fragments into the suicide vector pVA8912. Mutants defective in each of the individual genes (comC [SEQ ID NO:1], comD [SEQ ID NO:5], comE [SEQ ID NO:7]) are inactivated and their phenotypes are compared to the parent strain NG8 for their abilities to form biofilms, tolerate acidic pH (pH 2-4), and transport and incorporate DNA. The knockout mutants of com D [SEQ ID NO:5] and E [SEQ ID NO:7] were constructed by insertion-duplication mutagenesis, whereas the knockout comC [SEQ ID NO:1] mutant was created by allelic exchange via insertion of an erythromycin resistance determinant into the comC [SEQ ID NO:1] locus (Li et al, 2001). All mutant strains were therefore resistant to erythromycin. The wild-type strain was subcultured routinely on

Todd-Hewitt-Yeast Extract (THYE) agar plates (BBL®; Becton Dickinson, Cockeysville, MD), whereas the mutants were maintained on THYE agar plus 10 μg/ml of erythromycin. A minimal medium (DMM) was prepared to grow biofilms by a modification of the method described previously (Loo *et al*, 2000). The medium contained 58 mM K₂HPO₄, 15 mM KH₂PO₄, 10 mM (NH₄)₂SO₄, 35 mM NaCl, 2 mM MgSO₂·7H₂O, 0.2% (wt/vol) Casamino Acids and was supplemented with filter-sterilized vitamins, (0.04 mM nicotinic acid, 0.1 mM pyridoxine HCl, 0.01 mM pantothenic acid, 1 μM riboflavin, 0.3 μM thiamin HCl, and 0.05 μM D-biotin), amino acids (4 mM L-glutamic acid, 1 mM L-arginine HCl, 1.3 mM L-cysteine HCl, and 0.1 mM L-tryptophan) and 20 mM glucose.

[0090] To determine if the synthetic peptide [SEQ ID NO:16] could restore defective phenotypes of the comC [SEQ ID NO:2] mutants, a chemically synthesized 21-amino acid competence-stimulating peptide (CSP) [SEQ ID NO:4] (Li et al, 2001) was used in complementary experiments. The peptide was freshly dissolved in sterile distilled water to a concentration of 1 mg/ml. The CSP solution was then added to the cultures at a final concentration of 2 μg/ml 2 h after inoculation of bacterial cells.

[0096] Two methods modified based on the protocols described by Perry et al (Infect Immun, 41:722-727) and Lindler and Macrina (J Bacteriol, 166:658-665) were used to assay natural transformation of biofilm cells. Biofilms formed on polystyrene microtiter plates were added with 2 ml of pre-warmed, fresh THYE plus 5% horse serum (THYE-HS) immediately following removal of the BM medium, and the incubation continued at 37°C. After 2h incubation, a final concentration of 1 µg/ml plasmid DNA or 10 µg/ml of chromosomal DNA was added to each well. The cultures were incubated for an additional 2 h before collection of the cells for plating. To assay induction of competence by synthetic competence stimulating peptide (SCSP) [SEQ ID NO:16], the cultures were incubated for 30 min and a final concentration of 500 ng/ml of SCSP) [SEQ ID NO:16] was added to each well. After a 30 min incubation, equal amounts of DNA was added to each well (1 μg/ml plasmid or 10 μg/ml of chromosomal DNA) and incubation continued for another 2 h. Fluid medium was then removed from individual wells and the wells were washed once with PBS buffer. Biofilm cells were collected into 2 ml PBS buffer by gentle sonication or by washing the wells using a pipette. The samples were centrifuged at 12,000 X g for 5 min. Both biofilm and planktonic cells were resuspended into 200 μl of fresh medium and were immediately spread on THYE agar plates plus appropriate antibiotics. For the biofilms developed in the chemostat, rods with biofilm cells were removed and placed into 2 ml of pre-warmed, fresh THYE-HS medium for 30 min incubation. Transformation was then

initiated by using the same methods as described above. The planktonic cells were also removed to compare the transformation frequency. After completion of the transformation procedures, both biofilm and planktonic cells were spread on THYE agar plus appropriate antibiotic. Transformation frequency was assessed after 48-h incubation. Transformation frequency was expressed as the number of transformants per μg DNA per viable recipient at the time-of-DNA-added.

In the Claims:

- 3. (Amended) The compound of claim 2, comprising a derivative of [SEQ ID NO:2] or [SEQ ID NO:4], a fragment of [SEQ ID NO:2] or [SEQ ID NO:4] or a derivative of a fragment of [SEQ ID NO:2] or [SEQ ID NO:4].
- 4. (Amended) The compound of claim 3, wherein amino acids are removed from the N-terminus and/or C-terminus of [SEQ ID NO:2] or [SEQ ID NO:4].
- 5. (Amended) A pharmaceutical composition comprising all or part of the peptide of claims 1 to 4 claim 1 and a carrier.
- 6. (Amended) A method of medical treatment or prophylaxis of caries or endocarditis, comprising administering the compound of <u>claim 1</u> any of claims 1 to 4 or the pharmaceutical composition of claim 5.
- 7. CANCELLED
- 8. CANCELLED
- 9. CANCELLED
- 10. CANCELLED
- 11. CANCELLED
- 12. CANCELLED
- 13. CANCELLED
- 14. CANCELLED
- 15. CANCELLED
- 16. CANCELLED
- 17. CANCELLED
- 18. CANCELLED
- 19. CANCELLED
- 20. CANCELLED
- 21. CANCELLED

- 24. (Amended) The polypeptide of claim 23 comprising all or part of an amino acid sequence in [SEQ ID NO:2] or [SEQ ID NO:4].
- 29. (Amended) An isolated nucleic acid molecule encoding the polypeptide of any of claims 24 22 to 28.
- 30. (Amended) An antibody directed against the polypeptide of any of claims 21 22 to 28.
- 32. -- (Amended) A vaccine composition comprising all or part of the peptide of any of claims 21 22 to 28 and a carrier.
- 34. CANCELLED
- 35. CANCELLED
- 36. CANCELLED
- 37. CANCELLED

SEP 2 7 2001 SEP 2 7 2001 CON

APPENDIX - 2

COMPLETE SET OF PENDING CLAIMS, AS AMENDED ON SEPTEMBER 17, 2001

- 1. A compound that competitively inhibits binding of CSP to S. *mutans* histidine kinase.
- 2. The compound of claim 1, comprising a peptide or an antibody.
- 3. (Amended) The compound of claim 2, comprising a derivative of [SEQ ID NO:2] or [SEQ ID NO:4], a fragment of [SEQ ID NO:2] or [SEQ ID NO:4] or a derivative of a fragment of [SEQ ID NO:2] or [SEQ ID NO:4].
- 4. (Amended) The compound of claim 3, wherein amino acids are removed from the N-terminus and/or C-terminus of [SEQ ID NO:2] or [SEQ ID NO:4].
- 5. (Amended) A pharmaceutical composition comprising all or part of the peptide of claim 1 and a carrier.
- 6. (Amended) A method of medical treatment or prophylaxis of caries or endocarditis, comprising administering the compound of claim 1.
- 22. An isolated CSP or a fragment thereof having S. mutans CSP activity.
- 23. The polypeptide of claim 22 comprising a *S. mutans* CSP.
- 24. (Amended) The polypeptide of claim 23 comprising all or part of an amino acid sequence in [SEQ ID NO:2] or [SEQ ID NO:4].
- 25. A polypeptide fragment of the peptide of claim 24, or a peptide mimetic of the CSP.
- 26. The polypeptide of claim 24 which is recombinantly produced.
- 27. A polypeptide comprising a sequence having greater than 30%, 50% or 60% sequence identity to the polypeptide of claim 24.
- 28. The polypeptide of claim 24, isolated from S. mutans.
- 29. (Amended) An isolated nucleic acid molecule encoding the polypeptide of any of claims 22 to 28.
- 30. (Amended) An antibody directed against the polypeptide of any of claims 22 to 28.
- 31. The antibody of claim 30, comprising a monoclonal antibody or a polyclonal antibody.
- 32. (Amended) A vaccine composition comprising all or part of the peptide of any of claims 22 to 28 and a carrier.
- 33. The vaccine composition of claim 32, wherein the peptide is coupled to a compound comprising-all or part of KLH, ovalbumin, or thyroglobulin.

APPENDIX - 3 DETAILS OF DRAWING REVISIONS SUBMITTED ON SEPTEMBER 17, 2001

Appendix - 3



Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al

Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000 - Sheet 1 of 19

Figure 1

Schematic layout of the arrangement of the genetic locus encoding the signal peptide precursor, the histidine kinase and the response regulator. Note that this arrangement is different from other loci in related streptococci for the following reasons: a) The comegene is transcribed from its own promoter alone, unlike the genes thus far described in other streptococci that are arranged in an operon-like cluster with the come/DE genes being transcribed from a single promoter.

b) The comC gene is separated from the comD gene by 148 nuclectides.

Streptococcus mutans ComCDE Operon



Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al Atty Dkt No.: 1889-00401 Contact: C.M. Mintz (713)238-8000

Sheet 2 of 19

Figure 2

— Sequences of the open reading frames encoding the signal peptide precursor (ComC), the histidine kinase (ComD), and the response regulator (ComE).

S. mutans come gene

Encodes a precursor to a signal peptide
[ATGAAAAAACACTATCATTAAAAAATGACTTTAAAGAAATTAAGACTGATGAATTAG
AGATTATCATTGGCGGA (AGCGGAAGCCTATCAACATTTTTCCGGCTGTTTAACAGAAG
TTTTACACAAGCTTTGGGAAAA)]TAA

S. mutans CSP encoding sequence.

Competence Signal Peptide

AGCGGAAGCCTATCAACATTTTTCCGGCTGTTTAACAGAAGTTTTACACAAGCTTTGGG

AAAA [SEQ ID NO:1]

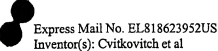
S. mutans comb gene

Encodes a protein that functions as a histidine kinase receptor

[ATGAATGAAGCCTTAATGATACTTTCAAATGGTTTATTAACTTATCTAACCGTTCTAT TTCTCTTGTTTCTATTTTCTAAGGTAAGTAATGTCACTTTATCGAAAAAGGAATTAACT CTTTTTTCGATAAGCAATTTTCTGATAATGATTGCTGTTACGATGGTGAACGTAAACCT GTTTTATCCTGCAGAGCCTCTTTATTTTATAGCTTTATCAATTTATCTTAATAGACAGA ATAGTCTTTCTCTAAATATATTTTATGGTCTGCTGCCTGTTGCCAGTTCTGACTTGTTT AGGCGGGCAATCATATTCTTTATCTTGGATGGAACTCAAGGAATTGTAATGGGCAGTAG CATTATAACCACCTATATGATCGAGTTTGCAGGAATAGCGCTAAGTTACCTCTTTCTCA GTGTGTTCAATGTTGATATTGGTCGACTTAAAGATAGTTTGACCAAGATGAAGGTCAAA AAACGCTTGATTCCAATGAATATTACTATGCTTCTATACTACCTTTTAATACAGGTATT GTATGTTATAGAGAGTTATAATGTGATACCGACTTTAAAATTTCGTAAATTTGTCGTTA TTGTCTATCTTATTTTATTTTTGATTCTGATCTCATTTTTAAGCCAATATACCAAACAA AAGGTTCAAAATGAGATAATGGCACAAAAGGAAGCTCAGATTCGAAATATCACCCAGTA TAGTCAGCAAATAGAATCTCTTTACAAGGATATTCGAAGTTTCCGCCATGATTATCTGA ATATTTTAACTAGCCTCAGATTAGGCATTGAAAATAAAGATTTAGCTAGTATTGAAAAG ATTTACCATCAAATCTTAGAAAAAACAGGACATCAATTGCAGGATACCCGTTATAATAT CGGCCATCTAGCTAATATTCAAAACGATGCTGTCAAGGGTATCTTGTCAGCAAAAATCT TAGAAGCTCAGAATAAAAAGATTGCTGTCAATGTAGAAGTCTCAAGTAAAATACAACTG CCTGAGATGGAGTTGCTTGATTTCATTACCATACTTTCTATCTTGTGTGATAATGCCAT TGAGGCTGCTTTCGAATCATTAAATCCTGAAATTCAGTTAGCCTTTTTTAAGAAAAATG GCAGTATAGTCTTTATCATTCAGAATTCCACCAAAGAAAAACAAATAGATGTGAGTAAA ATTTTTAAAGAAAACTATTCCACTAAAGGCTCCAATCGCGGTATTGGTTTAGCAAAGGT TATTCAAGCAACTCCTAATAATAAAA) TAG

S. mutans come gene

- Encodes a response regulator that activates transcription of a number of genes



Atty Dkt No.: 1889-00401 Contact: C.M. Mintz (713)238-8000

Sheet 3 of 19

Figure 2 (cont'd)

[ATGATTTCTATTTTTGTATTGGAAGATGATTTTTTACAACAAGGACGTCTTGAAACCA
CCATTGCAGCTATCATGAAAGAAAAAAATTGGTCTTATAAAGAATTGACTATTTTTTGGA
AAACCACAACAACTTATTGACGCTATCCCTGAAAAGGGCCAATCACCAGATTTTCTTTTT
GGATATTGAAATCAAAAAAAGAGGAAAAGAAAGGACTGGAAGTAGCCAATCAGATTAGAC
AGCATAATCCTAGTGCAGTTATTGTCTTTTGTCACGACACACTTCTGAGTTTATGCCCCTC
ACTTTTCAGTATCAGGTATCTGCTTTTGGATTTATTGAAACAGCCAGAAGAATG
GTCAATCAGAGGAACTTTTTATTTTCCATTCATCTGAAACTCAGGTCCCTTTT
GCTGAGATTCTGTATTTTGAAACATCTTCAACAGCCCATAAGCTCTGCCTTTATACTTA
TGATGAACGGATTGAATTCTACGGCAGTATGACATTGTTAAAATGGATAAGAGAC
TTTTTCAGTGCCATCGCTCTTTTATTTTTCCAATCCTGCCAATATTACCCGTATTGATCGG
AAAAAACGCTTGGCCTATTTTCGAAAATAATAAGTCTTGTCTTATTTCACGAACTAAGTT
AAAAAACGCTTGGCCTATTTTCGAAAATAATAAGTCTTGTCTTATTTCACGAACTAAGTT
AACAAAACTGAGAGCTGTGATTGCTGATCAAAGGAGAGACAAAA] TGA

Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000

Sheet 4 of 19

Figure 3

The amino acid sequences of the signal peptide precursor (ComC), the histidine kinase (ComD), and the response regulator (ComE).

S. mutans ComC protein (CSP Procursor)
MKKTLSLKNDFKEIKTDELEIIIGGSGSLSTFFRLFNRSFTQALGK

S. mutans comb protein (Histidine Kinase)

MNEALMILSNGLLTYLTVLFLLFLFSKVSNVTLSKKELTLFSISNFLIMIAVTMVNVNL

FYPAEPLYFIALSIYLNRQNSLSLNIFYGLLPVASSDLFRRAIIFFILDGTQGIVMGSS

IITTYMIEFAGIALSYLFLSVFNVDIGRLKDSLTKMKVKKRLIPMNITMLLYYLLIQVL

YVIESYNVIPTLKFRKFVVIVYLILFLILISFLSQYTKQKVQNEIMAQKEAQIRNITQY

SQQIESLYKDIRSFRHDYLNILTSLRLGIENKDLASIEKIYHQILEKTGHQLQDTRYNI

GHLANIQNDAVKGILSAKILEAQNKKIAVNVEVSSKIQLPEMELLDFITILSILCDNAI

EAAFESLNPEIQLAFFKKNGSIVFIIQNSTKEKQIDVSKIFKENYSTKGSNRGIGLAKV

NHILEHYPKTSLQTSNHHHLFKQLLIIK

S: mutans Come protein (Response Regulator)
MISIFVLEDDFLQQGRLETTIAAIMKEKNWSYKELTIFGKPQQLIDAIPEKGNHQIFFL
DIEIKKEEKKGLEVANQIRQHNPSAVIVFVTTHSEFMPLTFQYQVSALDFIDKSLNPEE
FSHRIESALYYAMENSQKNGQSEELFIFHSSETQFQVPFAEILYFBTSSTAHKLCLYTY
DERIEFYGSMTDIVKMDKRLFQCHRSFIVNPANITRIDRKKRLAYFRNNKSCLISRTKL
TKLRAVIADQRRAK



Express Mail No. EL818623952US

Inventor(s): Cvitkovitch et al Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000

Sheet 5 of 19

Figure 4

The deduced amino acid sequence of the signal peptide precursor in various strains and its predicted cleavage site. The original peptide is expressed as a 46 amino acid peptide that is cleaved after the glycine-glycine residues to generate an active signal peptide.

BM71 CSP	1 mkktpslkndfkeiktdeleiiiggsgslstffrlfnrsftqalgk 45
GB14 CSP	1 MKKTLSLKNDFKEIKTDELEIIIGGSGSLSTFFRLFNRSFTQALGK 46
H7 CSP	1 MKKTLSLKNDFREIKTDELEIIIGGSGSLSTFFRLFNRSFTOALGK 46
JH1005 CSP	1 MKKTLSLKNDFKEIKIDELEIIIGGSGTLSTFFRLFNRSFTQA 43
LT11 CSP	1 MKKTLSLKNDFREIKTDELBIIIGGSGSLSTFFRLFNRSFTQALGK 46
NG8 CSP	1 MKKTLSLKNDFKEIKTDELEITIGGSGSLSTFFRLFNRSFTQALGK 46
UAB159 CSP	1 MKKTLSLKNOFKEIKTDELEIIIGGSGSLSTFFRLFNRSFTQALGK 46
	**** ***********

consensus: 1 MKKTLSLKNDFKEIKTDELEIIIGG SGSLSTFFRLFNRSFTQALGK 46 predicted cleavage site:

Figure 5

The synthetic signal peptide that is effective at inducing competence, biofilm formation and acid tolerance in Streptococcus mutans.

SGSLSTFFRLFMRSFTQALGK [SEQ ID NO:2]

Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al

Atty Dkt No.: 1889-00401

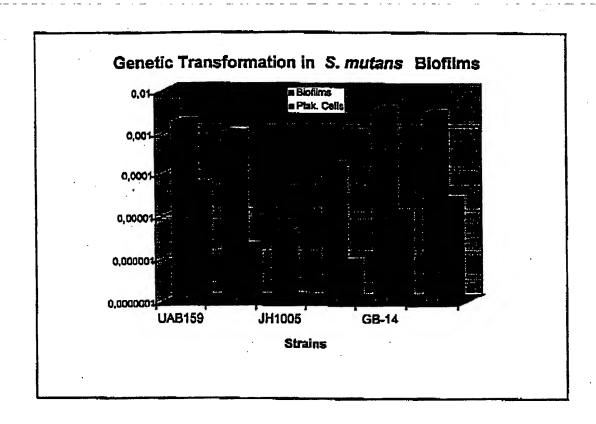
Contact: C.M. Mintz (713)238-8000 ---

Sheet 6 of 19

Figure 6

The natural activity of the signal/receptor system functioning in vitro in model biofilms
as determined by the ability of various strains of S. mutans to accept donor plasmid DNA

conferring crythromycin resistance.





Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000 -

Sheet 7 of 19

Figure 7

Induction of genetic transformation in Streptococcus mutans

by synthetic competence stimulating peptide (SCSP)

Strain	Peptide added Number of Transformants/Recipients	No peptide Number of Transformants/Recipients
UAB15	4.65 X 10 ⁻¹	1.78 X 10 ⁻⁶
JH1005 ²	6.98 X 10 ⁻²	0
hrrt 8		

The final concentration of SCSP used was 500 ng/ml.

The strain contains a nonsense mutation in the comC gene encoding the CSP.

express Mail No. EL818623952US

Inventor(s): Cvitkovitch et al Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000

Sheet 8 of 19

Figure 8

List of the primers used to amplify the genes or internal regions of the target genes by polymerase chain reaction (PCR) for subsequent sequencing or inactivation.

ComC region

ComC Primer Pair: F5-B5

[F5] 23406-23424 5'- AGTTTTTTGTCTGGCTGCG -3'
19 nt forward primer
pet G+C: 47.4 Tm: 50.5

[B5] 24056-24037 5'- TCCACTAAAGGCTCCAATCG -3' 20 nt backward primer pct G+C: 50.0 Tm: 51.9

651 nt product for F5-B5 pair (23406-24056) Optimal annealing temp: 50.3 pct G+C: 30.9 Tm: 71.5

ComD region

ComD Primer Pair: FI-B1

[F1] 392-415 5'- CGCTAAGTTACCTCTTTCTCAGTG -3'
24 nt forward primer
pct G+C: 45.8 Tm: 51.6

[B1] 683-663 5'- GCTTCCTTTTGTGCCATTATC -3'
21 nt backward primer
pct G+C: 42.9 Tm: 50.8

292 nt product for F1-B1 pair (392-683) Optimal annealing temp: 49.5 pct G+C: 30.8 Tm: 70.2

ComE region

ComE Primer Pair: F1-B1

[F1] 145-165 5'- CCTGAAAAGGGCAATCACCAG -3'
21 nt forward primer
pct G+C: 52.4 Tm: 55.9

[B1] 606-585 5'- GCGATGGCACTGAAAAAGTCTC -3'
22 nt backward primer
pct G+C: 50.0 Tru: 55.4

462 nt product for F1-B1 pair (145-606) Optimal annealing temp: 53.6 pct G+C: 38.3 Tm: 74.1 Express Mail No. EL818623952US

Inventor(s): Cvitkovitch et al Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000

Sheet 9 of 19

Figure 9	-
ComCDE local region. The ComC (first highlighted region; nucleotides 101 to 24	1),—
ComD (second highlighted region; nucleotides 383 to 1708) and ComE (third high	ilighted -
region; nucleotides 1705 to 2457) proteins are highlighted.	-

Sequence Range: 1 to 2557

10 20 30 40 50
ACATTATGTGTCCTAAGGAAAATATTACTTTTTCAAGAAAATCCATGATT
TGTAATACACAGGATTCCTTTTATAATGAAAAAGTTCTTTTAGGTACTAA
<K K L F I W S K

TITTCATARARAATAGTATACTARTTATAATCARARAAAAAGGAGATATAAAAAAAGGAGTATTTTTATCATATAGATTAATATTAGTTTTTTTCCTCTATATTTCKK M F F L I S I I I L F L L Y L

110 120 130 140 150
ATGAAAARACACTATCATTAAAAAATGACTTTAAAGAAATTAAGACTGA
TACTTTTTTTGTGATAGTAATTTTTTTACTGAAATTTCTTTAATTCTGACT

ORF RF[2]

160 170 180 190 200
TGAATTAGAGATTATCATTGGGGGAAGCGGAAGCCTATCAACATTTTTCC
ACTTAATCTCTAATAGTAACCGCCTTCGCCTTCGGATAGTTGTAAAAAGG
BTGTELTITTGGGGAAGCGSAAGCTTGTAAAAAGG

ORF RF[2]

<S N S I I M

210 220 230 240 250
GGCTGTTTAACAGAAGTTTTACACAAGCTTTGGGAAAATAAGATAGGCTA
CCGACAAATTGTCTTCAAAATGTGTTCGAAACCCTTTTATTCTATCCGAT
ET TO REF [2] >

260 270 280 290 300
ACATTGGAATAAAACAAGGCTGGATTTATTATTCCAGCCTTTTTAAATGT
TGTAACCTTATTTTGTTCCGACCTAATAATAAGGTCGGAAAAATTTACA

310 320 330 340 350
AAAATAAAAATACAGGGTTAAATAATCAAGTGTGCTGTCGTGGATGAGAA
TTTTATTTTTATGTCCCAATTTATTAGTTCACACGACAGCACCTACTCTT

360 370 380 390 400
GATAAAACTATCTCTTAGAGAATAGGCCTCCTCTATFTTATTATTAGGAG
CTATTTTGATAGAGAATCTCTTATCCGGAGGAGATAAAATAATAATCCTC

ek T 1 1 L CRF RF[Express Mail No. EL818623952US
Inventor(s): Cvitkovitch et al
Atty Dkt No.: 1889-00401
Contact: C.M. Mintz (713)238=8000

Sheet 10 of 19

Figure 9 (cont'd-1)

				. .		ORF						~~ <i>~</i> / <i>~</i> /			
									• •						
		460			47		~		80	3 ~~		490		n me	500 777
															GGA(
			I	H	M	V V		A	L		I	G	R	N	S
<h< th=""><th>B</th><th>L</th><th>_</th><th>-11</th><th>М</th><th>ORF</th><th></th><th></th><th>_</th><th>_</th><th>-</th><th>G</th><th>ж.</th><th>74</th><th>3</th></h<>	B	L	_	-11	М	ORF			_	_	-	G	ж.	74	3
						-									
	_	510				10						540			550
															GIT
		ACC													CAAI
	<u> </u>	حربين.		3.4	AL	ORF	RF	[4]	C	icrus.			W. YEE		er areas
						•									
		560			_	70						590			600
															TTA
AG	AAC	CAC	CII	AAG	AC	TAC	TAT	TTC	TGP	TAT	GA(GGI	AA	AAG	AAT
E	K.	En.	Ş	N	Q.";					E	5	G*	'n.		K
						ORF	r.E	[4]	C						
		610			62	20		6	30			640)		65
AAZ	AGGG	TAA		AAT	-I-I.(CAGG	ATT	TAA	TGF	TTC	:GAJ	VAGC	AGC	CTC	AAT
TT	rÇÇ	BATT	GAC	TIA	AA	STCC	AAT	ATT	ACT	CAAC	CT	FTÇG	TCC	GAC	TTA
															M
S.F		, T.	,			∴ಿ≣್			,,,,,,,	****					0.00
						ORF	RF				~ VAL				
					-	_		[4]	C_						
		660			-	- 70		[4]	C_ 80	-		€90			70
		TCAC	AÇ?		TĀ	- 70 GAAA	.GTA	[4]	C_ 80 TA	ATG)	laa:	690	LGC2	AACI	70 CCA
		TCAC AGTG	AÇ?		TĀ	- 70 GAAA	GTA CAI	[4] 6 TGC	C_ 80 TA	ATG)	laa:	690	LGC2	AACI	70 CCA AGGT
GI	AT	TCAC AGTG	AÇI TGT	TCI	ATA ATC	70 GAAA CTIT E OR	GTA CAI S LF R	[4] 6 TGG ACC M	C_ 80 TAI AT: V	atgi Pact M	LAA:	690 ICAA AGTI	AGCI ICGI S	AACI ITGA	70 CCA AGGT
GI	AT	TCAC AGTG	AÇI TGT	TCI	ATA AT	70 GAAA CTTT E OR	GTA CAI S IF R	[4] 6 TGG ACC M	C_ 80 TAI AT V	ATGI FACT M	AAX K	690 ICAA AGTI S	AGCI CG: S	AACI ITGA	70 CCA AGGT
GT	AT	TCAC AGTG	AÇI TGT	TCI	ATA AT	70 GAAA CTIT E OR	GTA CAI S IF R	[4] 6 TGG ACC M	C_ 80 TAI AT V	ATGI FACT M	AAX K	690 ICAA AGTI	AGCI CG: S	AACI ITGA	70 CCA AGGT
GT	AT	TCAC AGTG	ACI TGT H	TCI	TAC ATO	70 GAAA CTTT E OR	GTA CAI S IF R	[4] ACC M F[3]	C_ 80 TAI AT V	ATGI FACT M	AAX K	690 ICAA AGTI S	AGCI CG S	AACI ITGA	70 FCCA AGGT S
A	L X	TCAC AGTG S	ACI TGT H	K K	TAC I I	70 GAAA CTTT E OR OR	GTA CAI S IF R	[4] 6 TGG ACC M IF [3]	C BO TAI AT: V]	ATG! FACT M	AA K	690 FCAZ AGTT S	AGCI CG: S	AACI ITGA M	70 CCA GGT S
A	L	rcac AGTG S 000000 710	ACI TGI H	TCI K	TAC I I	70 GAAA CTTT E OR ORI 20 ACTT	GTA CAI S IF R	[4] 6ACC M F [3	C SO TO	ATG! FACT M	LAAT	690 ICAP AGTI S 740 GAC	AGC	AACI ITGA N E	70 CCA AGGT S
A	L AGG	rcac AGTG S 000000 710	ACI TGI H	TCI K	TAC I I	70 GAAA CTTT E OR OR OR ACTT	GTA CAT S LF R CTC	[4] 6ATGG ATGG M 1F [3] 7 [4]	C SO TO	ATG! FACT M	LAAT	690 ICAN AGTI S OGAL GAC	AGC	AACI ITGA N E	70 CCCA AGGT S
A A	L AGG	PCAC AGTG S 710 CAGT	ACZ TGT H	TATI	TAC I 7	70 GAAA CTTT E OR OR ACTT TGAA	GTA CAI S F R CAC CAC CAC CAC CAC CAC CAC CAC CAC CA	[4] 60 ACO M IF [3] FACTOR	C_ TAI TAI V I I C_ V30 TAG	ATG! M INCI IAC! IAC! T	AAA TTE K	690 ICAP AGTI S 740 GAC	AGC:	AACI ITGA M BATO TAO	70 FCCA GGT S
A A CTC	L AGG	PCAC AGTG S 710 CAGT	ACZ TGT H	TATI	TAC I 7	70 GAAA CTTT E OR ORI 20 ACTT TGAI	GTA CAT S IF F CGAO CTO E F F	[4] 66 ACC M F [3] F [4] F [4] T T T T T T T T T T T T T T T T T T T	C 80 AT AT C 30 C S S S S S S S S S S S S S S S S S S	ATGI M IACI IACI	AAA TTE K	690 ICAP AGTI S 740 GAC	AGC:	AACI ITGA M BATO TAO	70 CCCA AGGT S
A A CTC	L AGG	PCAC AGTG S 710 CAGT	ACZ TGT H	TATI	TAC I 7	70 GAAA CTTT E OR ORI 20 ACTT TGAI	GTA CAI S F R CAC CAC CAC CAC CAC CAC CAC CAC CAC CA	[4] 66 ACC M F [3] F [4] F [4] T T T T T T T T T T T T T T T T T T T	C 80 AT AT C 30 C S S S S S S S S S S S S S S S S S S	ATGI M IACI IACI	AAA TTE K	690 ICAP AGTI S 740 GAC	AGC:	AACI ITGA M BATO TAO	70 FCCA GGT S
A A CTC	L AGG	TCAC AGTG S 710 CAGTG GTCA S	ACI TGI H	TATI	TACE TO THE PERSON OF THE PERS	70 GAAA CTTT E OF ORI ACTT TGAI OI ORI	GTA CAT S IF F CGAO CTO E F F	[4] 66 CACO M F [3] F [4] F [4]	C S C S C C S C C C C C C C C C C C C C	ATGI M IACI IACI	AAA TTE K	690 AGTICAL S 740 GACICIGI	AGCI S S AGCI CGA	AACI ITGA M BATO TAO	70CA AGGT S
A CTC	AATI L RAGG AGG TCC G	710 CAGTG 710 CAGTG STCA	ACI TGT H TGT ACI	TTCI K	TACE TO THE TACE T	70 GAAA CTTT E OR OR 20 ACTTTGAA TGAA CR OR OR OR	GTA CAI S F F CGAC CGAC CGAC F F	[4] 66 ACC M IF [3] 7 [4] 7 [4] 7 [4]	C SO TAIL SO T	ATG:	AAA	690 ICAA AGTT S 740 GACI CTG T	AGCZ S S AGCZ A	AACI N N BATO	70CAAGGT S S TTTT AAAA F>
A CTC	AATI	710 CAGTG 710 CAGTG GTCA S	ACI TGT H TGT ACI	ETCI K	TAC	70 GAAA CTTT E OR OR 20 ACTTTGAA TGAA OR OR OR OR TTGAA TTGA	GTA CAT S IF F CGAC CTC S F F	[4] 66 ACC M 1F [3] 7 [4] 7 [4] 7 [4]	C SO	ATGI M IACI IACI T T	AAATTIAAATTAC	690 ICAA AGTT 740 GACI CTGT T	AGCI S S AAGCI CG A	AACTTGA N BATC	70CAAGGT S S S S S S S S S S S S S S S S S S
CTC CAC GAG S CEA	AATI L AGG	710 CAGTG 710 CAGTG GTCA S	ACI TGT H TGT ACI	ETCI K	TAC	70 GAAA CTTT E OR OR 20 ACTTTGAA TGAA OR OR OR OR TTGAA TTGA	GTA CAT S IF F CGAC CTC S F F	[4] 66 ACC M 1F [3] 7 [4] 7 [4] 7 [4]	C SO	ATGI M IACI IACI T T	AAATTIAAATTAC	690 ICAA AGTT 740 GACI CTGT T	AGCI S S AAGCI CG A	AACTTGA N BATC	70CAAGGT S S S S S S S S S S S S S S S S S S
CTC CAC SAC SAC TATA	AATI L RAGG AGG G TCT AGA F>	710 CAGTG TOCAGTG TOCAGTG TOCAGTG	ACI TGT H TGT ACI	ETCI K	TAC	70 GAAA CTTT E OR OR 20 ACTTTGAA TGAA OR OR OR OR TTGAA TTGA	GTA CAT S IF F CGAC CTC S F F	[4] 66 ACC M 1F [3] 7 [4] 7 [4] 7 [4]	C SO	ATGI M IACI IACI T T	AAATTIAAATTAC	690 ICAA AGTT 740 GACI CTGT T	AGCI S S AAGCI CG A	AACTTGA N BATC	70CA CCCA S S
CTC CAC S S S TAT ATA	ACG G TCT ACA TCT ACA F>	710 CAGTG TOCAGTG TOCAGTG TOCAGTG	ACI TGT H ACI C	TATA	TACARA AGA	70 GAAA CTTT E OR OR 20 ACTTTGAI TGAI OR OR 70	GTA CAI SF F CGAC CTC SF F CTC CAC CTC	[4] 66 ATGG ATGG ATGG ATGG ATGG ATGG ATGG A	C SO	ATGI M IACI IACI T T	ATTO	690 ICAA S 740 GACI CTG T	AGCI S AGCI AGCI A	AACITGA N N RATO I CAGO	70CAAGGT S S S S S S S S S S S S S S S S S S
CTC CAC S S S TAT ATA	ACG G TCT ACA TCT ACA F>	710 CAGTG S 710 CAGTG GTCA S 760 CAGGC	ACI TGT H ACI C	TATA	TACARA AGA	70 GAAA OF OR 20 ACTM L OR 70 TTGAAA	GTA CAI SF F CGAC CTC SF F CTC CAC CTC	[4] 66 KTGG KTGG KTGG KTGG KTGG KTGG KTGG K	C SO STANDARD C	ATGI M IACI IACI T T AGA	ATTO	690 ICAA S 740 GACI CTG T	AGCI S AGCI AGCI A	AACITGA N N RATO I CAGO	70CAAGGT S S S S S S S S S S S S S S S S S S
CTC CAG S S S TAT ATA	ACG G TCT ACA TCT ACA F>	710 CAGTG S 710 CAGTG GTCA S 760 CAGGC	ACI TGT H ACI C	TATA	TACARAGA	70 GAAA OF OR 20 ACTM L OR 70 TTGAAA	GTA CAI CAI F F PGA COTO E F F F F F F F F F F F F F F F F F F F	[4] 66 KTGG KTGG KTGG KTGG KTGG KTGG KTGG K	C SO STANDARD C	ATGI M IACI IACI T T AGA	ATTO	690 ICAA S 740 GACI CTG T	AGCI S AGCI AGCI A	AACITGA N N RATO I CAGO	70CAAGGT S S S S S S S S S S S S S S S S S S

Express Mail No. EL818623952US
Inventor(s): Cvitkovitch et al
Atty Dkt No.: 1889-00401
Contact: C.M. Mintz (713)238-8000
Sheet 11 of 19

Figure 9 (cont/d-2)

					°	••	,	3	<u> </u>						
	94	60		1	B70			88	0		8:	90		9	900
TGTC			واحتاسا			-1-1	רבטי		-	יייטיי	וינייניים	יממי	TACT	ישפני	7
										AGA					
ACAG			na.												~
H./: (C		X	**		Ţ							5.5.4.			+14
					O	C.F	RF	[4]	C						
	9	10			920			93	۸		q.	40		,) 50
ATCI			ም/3s			127	יכיים		_	الملت	_		ייין ייין	-	
TAGE															
TAG				4300				(77.74	No. Tier				47.61		;
SED IS		N. T.			0	RF	RF	[4]	C_	auer -			. Cook	75.3	
											_				
		60			970			98		<u></u> .	_	90			001
CATO															
GTA	CGC	CTI	TGA	AGC			_								
M	A	E	T	5	N	I	L	V	K	R.	F	Y	L	L	T
	- -	5:38-2	· S	R	7	D:	ALK.	. ¥.	* L.	···S	1/12/2	· Table	O	ok:	%s
Desire (Constitution of the Constitution of th	(3 . YE)		: حور ،				RF		C		-9-2-11	.,	111 76 5.	Charles, a	<i></i>
								•	-						
	10	10		1	020			103	30.		10	40		1	05
TAC	rggg	TGA	TAI	TTC	GAA	TC	rga(GCT:	rcc1	TTT	GTG	CCA	TTA	TCT	CA
	ACCC														
	L G					-		_	? I						
-			•			-	-	•			_		-	_	
792.56	. 444-119"	348.74	7584			21.77	14:48		3031	c° to	ori saliki	37 5 2 S 4 1		3.4	2.05
$X^{(i,j)}$				1 R			- 000040			Y-17-32		S is I	1	1.12	
					o	RF	RP	[4]	c _						
				_											
		160			.070			10				90			10
	AACC					-									
LAAC	TTGG	LAA	YAC	LAAC	CAT	AT.	AAC	CCV	ATT.	TTA	CIC	TAC	TCT	TAG	ΤT
L	N	L	L	F	G	I	L	A>							
									>						
50%	v	K'	O,	K.	T	Ϋ́	:Ö2	35%	1		S.	Í.	100	I.	Ľ
200	. Address	A-120 6 1	, and	.72 3	· <u> </u>	· · · ·	-414	[4]	C	(334)		7=-1	V	FT 6.1.	
	, <u></u>							,	_						
	71	110		1	1120)		11	30		11	L 4 0		i	1.5
A POTA	LAAA		27.ID1				~~~			<u>የ</u> አ <i>ሶር</i>			מ מידיי		
	TTT						•								
TRAT		CT.T.	11.4		T. T.	T		***		17 MAG	10 E	LAM	**************************************	100	.,
	· ·	Ţ	· · · · · · ·	∴. :V:,	#									- <u>T</u>	٠. ١
						RP	RF	[4]	c _						
														_	
		160		_	1170			11				L90			.20
rate	ACA"	LTA:	CAA	CTCI	CTA	TA	ACA	TAC	AAT.	ACC	'GT)TT	AAAZ	\GGT	AG
ATAG	TGT	LAT	VII.												
٠. تا >	y : 1	T	Ž · · ·	5 " I	• • •		V.	Y 4	L	V (Σ,	r vi	, F	, <u>.</u>	
2 .0. 20				•				[4]		• • •			, , ,,,,,	<i>,</i> , ,	
	_				`			1	-						
	12	210		2	L220)		12	30		1:	240		1	.25
ATAG			AGT	_			TGC	AAT	CAA	GCG:	LL.L.	TTT	SACO	TTC	Αī
TATO										•		_	-		
**************************************	27.25 A. 41			_ ,										** ** ****	
5			•	*:				[4]				1		X.	, MI

Express Mail No. EL818623952US
Inventor(s): Cvitkovitch et al
Atty Dkt No.: 1889-00401
Contact: C.M. Mintz (713)238-8000
Sheet 12 of 19

Figure 9 (cont

_					,							`					
		260			127				280			12				1	3 (
TTGG AACC	T CAL	AAC TTC	ייייען בייייען	CII	Z TAM	rea.	CGA	CCA	ATA Ta	TTCA	MC!	**************************************	iA.	ACZ	CA	CIC	32
	V. 55			1			5-C1		Ť		4627	AH	7	, O		GA.	
Apple 1 (1985)				e e	* #		PR						Z E	2001	Carrette.	3.1	
						<u>-</u> 							_	_	-	- 0	
AAAG		310 TAA		'AGC	132 GC		TCC'		330 222		Y23.	134 222			·	13	3 :
TTTC	TCC	TTA	GAA	TCG	CGI	YΑ	AGG	ACG	III	GAG	CT	r Carry	Les. Ler	LANC LANC	ירטי ינטי	3G.	
218	L.	Ý	SU	17.	À	1	G	A		E		76 N	3	74.5	77	Ť	÷
					-		PR										ردد
	1.	360			137	70		1	380	1		139	20			٠,	
TAAT						_	rrc				ימי:			יי מיבי די מיבי	ממי	14 8/27	.,
ATTA	CGA:	TGA	CGG	GTA	ATC	TT.	AAG	CAA	CTC	AAG	GT	LGGT	املما	ייייי בידי	املاملاء محمد		<u>~</u>
	S	ş	Ğ	M	v	I,	,G	∵ ହ	4.1		36.7			Œ,		- 1	
						ORI L	FR E	F [4] K] C		M	W	_	5	ь	s	
					<_							•••			~		
	14	110			142	0		1.	430			144	L Ch			14	
ATGA:							rca(ACI			ECA	GAC	CZ	v.
TACT	AAÇ	3GG	CGG	ATT	TGI	TC	GT	CIT	GAC	CGI	TG:	rcçe	T	GI	CIC	3G1	
	A	PR	R		*/I	ORI		S. C.	S.V.	$\mathbf{A}^{*}\lambda^{\dagger}$	Y N	PX	T,	, I	松		3
<\$	ō	G	G	L	C	T	r Ki	? [4] V	l C		Ī		•	С	v	P	_
AAAT?		460 TTT	AGA.		147		\\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-	14	180	A PROP	***	145	0			15	
AAAT? TTTA:	ATA:	rrr.	aga TCT	GAA	AGA	CT	TAAC	TG:	CT	ATT	AAC	ATA	Ā	TT	GA:	מבי	J
AAAT ATTA	ATA:	TT.	TCT	GAA CTT	aga TCT	CTA GAI	CAAC	erg: Saci	rct Aga	ATT AAT	TTC	ATA	A	CAA	CTZ	יי דידי	7
AAAT ATTT	ATA:	TT.	TCT	GAA CTT	aga TCT L	CTA GA1	CAAC	PTG:	ICT AGA R	ATT AAT	TTC	ATA	A	CAA	CTZ	יי דידי	7
AAAT TTTA:	ATA:	TT.	TCT	GAA CTT SIZ	aga TCT	CTA GAT S ORI	AAC N	PACI Q	TCT AGA R	TAA N	TTC	TATA	A	CAA	CTZ	TAZ YTT	
TTTA	ATA: PATZ Fail	TT.	ict E	GAA CTT S	AGA TCT L	CTA GAT S ORI	AAC N	27G: 3AC: Q: 7 [4]	PCT AGA R C	TAA	Ti	TATA	AJ	PAA V	CTZ S	TAR ATT	
CTAT	ATAT PATA I I I I I	TT.	ict Light Aag	GAA CTT ST	AGA TCT 152 CTC	GAT S ORI O TGC	TAAC RI TAGC	TG: PACI P [4]	PCT AGA R C S 3 0 AAA	ATT TAA N	TTO	TATA TATA TATA TATA	AJ TI	AA?	CTZ	TAR ATT	
CTAT	ATAT PATA I I I I I	TT.	ict Light Aag	GAA CTT ST	AGA TCT 152 CTC	GAT S ORI O TGC	TAAC RI TAGC	TG: PACI P [4]	PCT AGA R C S 3 0 AAA	ATT TAA N	TTO	TATA TATA TATA TATA	AJ TI	AA?	CTZ	TAR ATT	
CTAT	ATAT PATA I I I I I	TT.	ict Light Aag	GAA CTT ST	AGA TCT 152 CTC	GAT GAT ORI O TGC	TAAC RI TAGC	TA	PCT AGA R C S S S S S S S S S S S S S S S S S S	ATT TAA N	TTO	TATA TATA TATA TATA	AJ TI	AA?	CTZ	TAR ATT	
CTAT	ATATI	TT.	ict Light Aag	GAA CTT ST	AGA TCT 152 CTC	GAT S ORI ORI ACO	AGC	TATA	R R R S S S S S S S S S S S S S S S S S	ATT TAA N ACA TGT	TTO	154	O CC	AA?	CTZ	15 CA	
CTATU GATA SI	ATATATATATATATATATATATATATATATATATATAT	SIO TATA	AAG ITC	GAA CTT ST? AGG TCC	152 CTC GAG	GAT GAT ORI ACC ACC GAA	AAC RI AGC TCC RI	ITG: SAC: Q: 1.1 SAT: TA: [4]	RGA RGA RGA RGG RGG RGG RGG RGG RGG RGG	ACA TGT	GGI	154 TAI 154 TTA TAAT 159	O AC	TAR TT	CAC	15 CA IS CA IS	
TATE TATE TATE TATE TATE TATE TATE TATE	ATATATATATATATATATATATATATATATATATATAT	SIO	AAG FTC	GAA CTT S AGG TCC TTA AAT	AGA TCT 152 CTC GAG 157 TCA	GATI GATI ORI ORI ORI ORI ORI ORI ORI ORI ORI OR	AGC TCC RI	TIGO	TCTAGA	ACA TGT	GGI	154 TAT 154 TTA AAT 159	O AG	TAR AG	CAC	1S CA SGI	
CTATI	ATATATATATATATATATATATATATATATATATATAT	SIO	AAG FTC	GAA CTT S AGG TCC TTA AAT	AGA TCT 152 CTC GAG B	GAN GAN ORI ORI ORI ORI GAN	AAC	TAC	TCT AGA E C 530 AAA FTT C 680 TT	ATT TAA ACA TGT ATC	GGI	154 TAT 154 TTA AAT 159	O AG	TAR AG	CAC	1S CA SGI	
CTATI	ATATATATATATATATATATATATATATATATATATAT	SIO	AAG FTC	GAA CTT S AGG TCC TTA AAT	AGA TCT 152 CTC GAG B	GAN GAN ORI ORI ORI ORI GAN	AGC TCC RI	TAC	TCT AGA E C 530 AAA FTT C 680 TT	ATT TAA ACA TGT ATC	GGI	154 TAT 154 TTA AAT 159	O AG	TAR AG	CAC	1S CA SGI	
CTATI GATA CTAR CATA	ATATA	S10 S10 S10 S10 S60 SA	AAG ITCA ICA AGT	GAA CTT AGG TCC P	AGA TCT 152 CTC GAG 157 TCA AGT 162	GAN ORI ORI ORI ORI ORI ORI ORI	IAAGO RE	PER CONTROL OF THE PER CONTROL O	TCT AGA R C S S S S S S S S S S S S S S S S S S	ACA ACA TGT	GGI GGI GAI	154 154 171 159 159 164	O AG	AG TO	CAC	15 CA 16 CA 16 TA	
CTATE GATA GTAAC CATTO	ATA: I AT	S10 NTA	AAG HTCA HCA AGT	GAA CTT SI AGG TCC P	152 CTC GAG 157 TCA AGT 162 ACA	GAACO CATA	IAAC IAGO ITCO IAAC ITTO IAAC ITTO ITTO	PER CONTRACTOR OF THE	AGA C C SAA C T T C C C C C C C C C C C C C C C C	ATT ACA TGT ATC	GGI GCL GAL GAL GAL GAL GAL GAL GAL GAL GAL GA	154 154 17TA 17TA 159 164 17TA	O COCO COCO COCO COCO COCO COCO COCO C	AG TO	CACGTO	15 CA 16 16 16 16	
CTATI GATA GTAAC CATTO	ATA: I AT	S10 NTA	AAG TCA TCA AGT MAAA	GAA CTT SI AGG TCC P	152 CTC GAG 157 TCA AGT 162 ACA	ORI ORI ORI ORI ORI ORI ORI ORI ORI ORI	AACO RE	PRODUCTION OF THE PRODUCTION O	TOTAGA CONTRACTOR CONT	ATT ACA TAGA ATC TAGA	GGI CCI CAI CTI	154 154 17TA 17TA 159 164 17TA	O COCO COCO COCO COCO COCO COCO COCO C	AG TO	CACGTO	15 CA 16 16 16 16	
CTATI GATA GTAAC CATTO	ATA: I AT	S10 NTA	AAG TCA TCA AGT MAAA	GAA CTT SI AGG TCC P	AGA TCT 152 CTC GAG 157 TCA AGT 162	ORI ORI ORI ORI ORI ORI ORI ORI ORI	IAAGO TOO TOO TOO TOO TOO TOO TOO TOO TOO T	TIGO LACO LACO LACO LACO LACO LACO LACO LAC	C SO C C C C C C C C C C C C C C C C C C	ATT ACA TAGA ATC TAGA	GGI CCI CAI CTI	154 154 17TA 17TA 159 164 17TA	O COCO COCO COCO COCO COCO COCO COCO C	AG TO	CACGTO	15 CA 16 16 16 16	
CTATI GATA GTAAC CATTO	ATA: I AT	S10 S10 S60 SAN S10 SAN S10	AAG PICA AGT AAAA	GAA CTT S S AGG TCC P TTA AAT CAC	152 162 162	GATE ORIGINAL ORIGINA	RAGO RECEIVED RECEIVE	TIGO LACO LACO LACO LACO LACO LACO LACO LAC	FOTTAGA STATE OF THE STATE OF T	ATT ACA TAGA ATC TAGA	GGI CCI CAI CTI	154 154 154 159 164 164 174 164	O C G C T C AATT	AG TO	CACGTO	15 CA 16 AAT 16 AAT 16 AC	
CTATI GATA CATAC CATTO	ATA: I AT	S10	AAG FICA AGT NAAA	GAA CTT ST AGG TCC P	AGA TCT 152 CTC GAG B 157 TCA AGT 162 ACA 167	GAMACTTI	RAGO RE	Property of the control of the contr	FCT AGA FT C SAA FT T	ATT TAA ACA ATC TAGA ACA ACA ACA ACA ACC TAGA ACA ACC TAGA ACC TAG	GGI CCI GAI GAI TTI	154 154 154 159 164 164 164	ATTEN O ACTION O ATTENDED	AC TG	CAC GTC GTC AAI	15 CA ATTA 16 ATTA 16 ATTA 17	
CTATI SATAC STRAC SATAC SATAC	ATA: I AT	S10	AAG HTCA HTCA AGT HAAA	GAA CTT ST AGG TCC P	152 162 167 167 167	GAM ORI	TAACO RECEIVED RECEIV	TG: Q:	TOTT AGA AGA AGA AGA AGA AGA AGA AGA AGA A	ACA TGT ACA ATC TGG TGG AGA ACA	GGI GCC GAL GAL GAL GAL GAL GAL GAL GAL GAL GAL	154 154 159 159 164 174 164 169	ATTEN OCCUPANT O ATTEN OCCUPANT OCCUPA	AC TG	CACGTO	1SGIMATA 16GAGTC	THE STATE OF ASS - SATE OF
CTATI GATA GTAAC CATTO	ATA: I AT	S10	AAG HICA HICA AGT HITCA HICA HICA HICA HICA HICA HICA HICA HI	GAA CTT S AGG TCC P TTA AATI CAC	152 162 167 167 167	GAN ORI ORI ORI ORI ORI ORI ORI ORI ORI ORI	TAACO RECTOR REC	1: [4] 1: [4] 1: [4] 1: [4] 1: [4] 1: [4] 1: [4] 1: [4] 1: [4] 1: [4]	FOTTAGA FOR THE STATE OF THE ST	ATT TAA ACA ATC TAGA ACA ACA ACA ACA ACA ACA ACA ACA ACA	GGI GCC GAP GAP GAP GAP GAP GAP GAP GAP GAP GAP	154 154 159 159 164 174 164 169	OCCUPANT OF ANTICE	AGC TA	CTAC STORY GTO TTA AAC TTC	1SCATION 160 TO COLUMN 170 CCC	

Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al Atty Dkt No.: 1889-00401 Contact: C.M. Mintz (713)238-8000 Sheet 13 of 19 Figure 9 (contd-4) 1730 1710 TCATTCATTTTGCTCTCCTTTGATCAGCAATCACAGCTCTCAGTTTTGTT agtaagtaaaacgaggaaactagtcgitagtgtcgagagtcaaaacaa E N M ALWAY ROOM OF A TOWN ASTR ORF RF[5] C 1790 AACTTAGTTCGTGAAATAAGACAAGACTTATTATTTCGAAAATAGGCCAA TTGAATCAAGCACTTTATTCTGTTCTGAATAATAAAGCTTTTATCCGGTT MAKET BUS TYD CUSOKYNON BUSETY AUT ORF RF[5] C 1810 1830 1840 CCCTTTTTTCCCATCAATACGCCTAATATTGGCAGGATTGACAATAAAAG CGCAAAAAAGGCTAGTTATGCCCATTATAACCGTCCTAACTGTTATTTTC THE RESIDENCE OF THE RESIDENCE OF SECTION SECT ORF RF[5] C 1880 1860 MALKKSLIHFNNVSH> ORF RF[1] ER H.C. O. P. L. R. K. D. M. K. V. I. D. T. M ORF RF[5] C CTCCCGTAGAATTCAATCCGTTCATCATAAGTATAAAGGCAGAGCTTATG GACGGCATCTTAAGTTAGGCAAGTAGTATTCATATTTCCGTCTCGAATAC TAVEFNPFIISIKAELM> ORF RF[1] L'REDYTOTERE ORF RF[5] C GNMML ORF RF[6] C 1970 1980 1990 **GCCTGTTGAAGATGTTTCAAAATACAGAATCTCAGCAAAAGGGACCTGAA** CCGACAACTTCTACAAAGTTTTATGTCTTAGAGTCGTTTTCCCTGGACTT G C> SAS TEFFERENCE POVED TE ORF RF[5] C

2010 2020 2030 2040 2050
ACTGAGTTTCAGATGAAATGAAAAAAAAAGTTCCTCTGATTGACCATTC
TGACTCAAAGTCTACCTTTATTTTTTCAAGGAGACTAACTGGTAAG
SO T E S S H F I B T E E S O G N
ORF RF[5] C

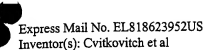
LICFR ORFRF[6] C

<S L R L H I S F L F N R Q N V M F

Express Mail No. EL818623952US
Inventor(s): Cvitkovitch et al
Atty Dkt No.: 1889-00401
Contact: C.M. Mintz (713)238-8000
Sheet 14 of 19

Figure 9 (confld-5)

	_	_	_		_				RF		_							_		
		-	60			20	70			208	'n			20	90				110	n
110/w	TV2/				TCC	ATA:		AT7	יייבו			CTY	נבב				CGC	_		-
						TAT														
75.	52	18,	N		3	M	Ã:	Y.	×Υ	Ľ	V A									•
	***	11.2.11				`				[5]					~~~			10.0		
<r< td=""><td></td><td>A OP F</td><td>T</td><td>X</td><td>-</td><td>C</td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></r<>		A OP F	T	X	-	C														
		~~~			-,															
		21	10			21	20			21:	30			23	L <b>4</b> 0	)		2	215	O
AA	CT	CCI	'CA	GG	AT.	CAZ	\AG	AT.	FIA	<b>ICA</b>	ATA	AA	AT(	c	W	\GC	AG	ATZ	ACC	T
						AGTT														
F.	J	77	72	ê.	N.	L	ZS.	. Pi	Ž.	D		F.	Ď			Αř	°S:		1	Q
							_0	RF	RF	[5]	C_									
						23													22(	
						AGGG														
TA	ΤG	ACI		TC	AC.	rcco	CG	TA.	1.1.I.	GAG	rCı									
										•		M	٠,	C	R	D	,	K	D:	>
37	2.5		: :-:::			್ಣಿಂ		w S	77	. E			10			300	1011	-	37	_
- T	bie.	Y		EIE-						[5]	2	NO TE				1500				
	_							~~~			_						_		_	_
		22	210	)		22	220			22	30			2	24(	)			22!	50
ATA	AC	TG	CAC	TA	GG	ATT	ATG	CT	GTC	TAR	T¢:	GA	TT	GG	CT	ACI	TC	CA	GT	CC
ΑŢ	TC	AC	3TG	ΑT	CC'	TAA:	[AC	GA	CAG	ATT	AG/	CT	AA	CC	GA:	rgz	AG	GT	CA	36
•	N	C	7	•	R	I	M	Ŀ	S	N	I		I	G	7	Y	F	Q		5>
r ·	W.	A	3	1	Ð.	, M.	H	Q	R	Ψī		2	ŊŶ.	A	en.	Ų, į	B	ت	1	3
							0	K.	KE	. [ 2 ]	_ C_									
			250							[5]	<u>د</u> 8۸	····			29				<b>-</b>	0.0
******	·		260		-141	2:	270	ì		22				_	29 nn	_			23 TG	
	-	TT	rcc	TC		2: TTT	270 TGA	TT	TCA	22 Ata	TC			AG	AA	Laa		GG	TG	ĽA
	\G/	TT AA	rcc	TC		2: TTT AAA	270 TGA	TT	TCA AGI	22 ATA TAT	TC	TT	TT	AG TÇ	AA	Laa		GG .CC	TG	ľA LT
VA.	\G/	TT AA	TCC AGC	TC AG	AA	2: TTT AAA	270 IGA ACI	TT	TCA	22 ATA TAT	TC( AG(			AG TÇ	AA TT	AAT TTI	AGA	GG .CC	TG	ľA LT
VA.	\G/	TT AA	TCC AGC	TC AG L	aa F	2: TTT AAA	270 IGA ACI	TT 'AA	TCA AGI F	22 ATA TAT N	TC( AG( I	Q Q	TT	AG	aa TT B	AAT TTI	AGA	GG .CC	TG	ľA VI
VA.	\G/	TT AA	TCC AGC	TC AG L	aa F	2: TTT AAA F	270 IGA ACI	TI	TCA AGI F	22 ATA TAT N	TCC AGC I	Q Q	TT	AG	aa TT B	AAT TTI	AGA	GG .CC	TG	ľA LT
VA.	\G/	TT AA	TCC AGC	TC AG L	aa F	Z: TTT AAA F	270 IGA ACT I	TT	TCA AGI F	ZZ ATA TAT N	TCC AGC I D	Q Q	TT	AG	aa TT B	AAT TTI	AGA	GG .CC	TG. V	LA LI L
F	G/	AA Q	TCC AGG F	AG L	P	2: TTT AAA F	270 FGA ACT D	TT	TCA AGI F	ZZ ATA TAT Y (5)	TCC AGC I D	C C	K	AG TC	TT E	TTI N N	I.	GG CC	TG AC V	AI IA I
F	G	AA 2	TCC AGG F 310	E C	AA F K	Z: TII AAA F K	270 IGA ACT D	AA PRF	TCA AGI F RI	ZZ ATA TAT N [5]	TCC AGC I D C 30	Q (L)	K	AG TC	TT E	N N	AGA	DD.	TG AC V	AI II ji
F F IGO ACO	G/ I	AA 2	TCC AGC F 31( TTC	E CAC	AA F	Z: TTT AAA F K Z ATA TAT	270 IGA ACT D 320 GCG	TT AA ) ) RF	TCA AGI F RI	ZZ ATAT TAT N [5]	TCC AGC I Do C	Q L	K	AG TC	AA TT B	N N N O TTO	AGA	GG CC	TG AC V Zi	TA 1 5 ( TA
F F IGO ACO	G	AA 2	TCC AGC F 31( TTC	E CAC	AA F	Z: TII AAA F K	270 IGA ACT D 320 GCG	AA PRF	TCA AGI F RI	ZZ ATA TAT N [5]	TCC AGC I Do C	Q L	K	AG TC	AA TT B	N N N O TTO	AGA	DD.	TG AC V Zi	TA 1 5 ( TA
F F IGO ACO	G/ I	AA 2	TCC AGC F 31( TTC	E CAC	AA F K GG	Z: AAA F K X X ATA TAT	270 IGA ACT D 320 GCG CGC	TT TAA PRF VTC	TCA AGI F RI AAI	ZZ ATA TAT N [5]	TCC AGC I D C TTC AA	Q L	K	AG TC 2 GG CC W	AA TT E 34 TT AA	AATTI N N N N N N N N N N N N N N N N N N	AGA L Se Q Se T	GG CC	TG AC V 23 AA TT	AI AI I SO AI AI
F F IGO ACO	G/ I	AA 2	TCC AGC F 31( TTC	E CAC	AA F K GG	Z: TTT AAA F K Z ATA TAT	270 IGA ACT D 320 GCC GCC	PAR V	TCA AGI F RI RI TTI	ZZ ATA TAT W [5] Z3 AAG TTC K	TCO AGO I D C 30 TTO AA	Q Q Z	K	AG TC 2 GG CC W	AA TT E 34 TT AA	AATTI N N N N N N N N N N N N N N N N N N	AGA L Se Q Se T	GG CC	TG AC V 23 AA TT	AI AI I SO AI AI
F F IGO ACO	G/ I	AA 2	TCC AGC F 31( TTC	E CAC	AA F K GG	Z: AAA F K X X ATA TAT	270 IGA ACT D 320 GCC GCC	PAR V	TCA AGI F RI RI TTI	ZZ ATA TAT N [5]	TCO AGO I D C 30 TTO AA	Q Q Z	K	AG TC 2 GG CC W	AA TT E 34 TT AA	AATTI N N N N N N N N N N N N N N N N N N	AGA L Se Q Se T	GG CC	TG AC V 23 AA TT	AI I I SO E AI AI
F F IGO ACO	G/ I	Z Z Z Z Z Z Z Z	TCC AGC F 31( TTC	ACCEPTAGE OF THE PARTY OF THE P	AA F K GG	2: TTT AAA F X X ATA TAT D	2700 TGAACT D	PAR PAR PAR V	TCA AGI F RI AAI N	ZZ ATA TAT W [5] Z3 AAG TTC K	TCC AGC I D G G TTC AA L	Q L	K	AG TC 2 GG CC W	AA TT E 34 TT AA	AAA	AGA L Se Q Se T	GG CC	TG AC V 23 AA TT	SO S
F CACC	IGI I I I I I I I	2 TT GAA L	TCC AGG F 31() TTC AAC F	L L L L L L L L L L L L L L L L L L L	F GGGCCC	2: TTT AAA F X X ATA TAT D	270 TGA ACT D 320 GCG CGC S	PRE	TCA AGI F RI AAII N	ZZZATA TAT W TAT (5)	TCC AGC I D TTC AA L C	Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q	K	2 GGG W	TT E 34 TT AA P	AAC C	L CA	AA TI	TG AC V 23 AA TI V	TATE OF THE PROPERTY OF THE PR
TGC ACC	CC GG	2 TI GAA L 2 ATT	TCC AGG F 31( TTC AAC F 36( CT	TO LACOUNT OF THE PARTY OF THE	AA F	2: TIII AAA F X AIA TAT D	270 TGA ACT D 320 GCC S A	PER V D CAN DE LA CENTRAL DE L	TCA AGT F RI AAT TTA	ZZZATAT TAT TAT TAT (5)	TCC ACC I D C 30 TTC AA L C 80 CT	Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q	K	AG TC 2 GG CC W	34 TT RAA P	AATI N O CTTO	IGA I GCA I GCA I GCA A GCA	AAATT	TG AC V Z3 AA TT V Z4 C	A)
F F F F F F F F F F F F F F F F F F F	CC GG	2 TI GAA L 2 ATT	TCC AGG F 31( TTC AAC F 36( CT	TO LACOUNT OF THE PARTY OF THE	AA F	2: TIT AAA F X X ATA TAT D	270 TGA ACT D 320 GCC S A	PER V D CAN DE LA CENTRAL DE L	TCA AGT F RI AAT TTA	ZZZATAT TAT TAT TAT (5)	TCC ACC I D C 30 TTC AA L C 80 CT	Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q	K	AG TC 2 GG CC W	34 TT RAA P	AATI N O CTTO	IGA I GCA I GCA I GCA A GCA	AAATT	TG AC V Z3 AA TT V Z4 C	ATATION OF COMMISSION OF COMMI
F F F F F F F F F F F F F F F F F F F	CC GG	2 TI GAA L 2 ATT	TCC AGG F 31( TTC AAC F 36( CT	TO LACOUNT OF THE PARTY OF THE	AA F	2: TIII AAA F X AIA TAT D	270 TGA ACT D 320 GCC S A	PER V D CAN DE LA CENTRAL DE L	TCA AGT F RI AAT TTA	ZZZATAT TAT TAT TAT (5)	TCC ACC I D C 30 TTC AA L C 80 CT	Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q	K K K K K K K K K K K K K K K K K K K	2 GG CC W	34 TT B 34 TT AA F	N O C C C C C C C C C C C C C C C C C C	CCA CCA CCA CCA CCA CCA CCA	AA TI K	TGAC V Z3 AAA TT N Z4 TGC	ATA TA 50 ATA OGG
TGC ACC	CC GG	2 TI GAA L 2 ATT	TCC AGG F 31( TTC AAC F 36( CT	TO LACOUNT OF THE PARTY OF THE	AA F	2: TIII AAA F X AIA TAT D	270 TGA ACT D 320 GCC S A	PER V D CAN DE LA CENTRAL DE L	TCA AGT F RI AAT TTA	ZZZATAT TAT TAT TAT (5)	TCC ACC I D C 30 TTC AA L C 80 CT	Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q	K K K K K K K K K K K K K K K K K K K	2 GG CC W	34 TT B 34 TT AA F	AATI N O CTTO	CCA CCA CCA CCA CCA CCA CCA	AA TI K	TG AC V 23 AA TI N V-I 24 VTC	ATA TA 50 ATA OGG



Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000

Sheet 15 of 19

Figure 9 (cont'd-6)

2410 2420 2430 2440 2450
GGTTTCAAGACGTCCTTGTTGTAAAAATCATCTTCCAATACAAAATAG
CCAAAGTTCTGCAGGAACAACATTTTTTAGTAGAAGGTTATGTTTTATC
G P K T S L L>

V S R R P C C K K S S S N T K I>

ORF RF[5] C

2460 2470 2480 2490 2500
ARATCATTATTTCTCCTTTAATCTTCTATTTAGGTTAGCTGATTAACACT
TTTAGTAATAAAGAGGAAATTAGAAGATAAATCCAATCGACTAATTGTGA
E I I S P L I F Y L G>

*L M

2510 2520 2530 2540 2550 ATACACAGAAAAGGTATAAAACGATATCACTCAATAAAATCTACTAACTT TATGTGTCTTTTCCATATTTTGCTATAGTGAGTTATTTTTAGATGATTGAA

AATAACC TTATTGG xpress Mail No. EL818623952US inventor(s): Cvitkovitch et al

Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000

Sheet 16 of 19

Figure 10

The comX nucleotid sequence, amino acid sequence, and its local region with 100bp included both upstream and downstream (promoter is upstream).

A.)
S. mutans comx gene

S. mutans ComX protein

MEEDFEIVFNKVKPIVWKLSRYYFIKMWTREDWQQEGMLILHQLLREHPELEEDDTKLY IYFKTRFSNYIKDVLRQQESQKRRFNRMSYEEVGEIEHCLSSGGMQLDEYILFRDSLLA YKQGLSTEKQELFERLVAGEHFLGRQSMLKDLRKKLSDFKEK

S. mutans comx gene local region-

Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al

Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000

Sheet 17 of 19

Figure 11.

The comA and comB nuclectide and amino acid sequences. ComA and ComB are the components of the CSP exporter.

S. mutans com/ gene

ATGAAACAAGTTATTTATGTTGTTTTAATCGTCATAGCCGTTAACATTCTCTTAGAGAT TATCAAAAGAGTAACAAAAAGGGGAGGGACAGTTTCGTCATCTAATCCTTTACCAGATG GGCAGTCTAAGTTGTTTTGGCGCAGACATTATAAGCTAGTACCTCAGATTGATACCAGA GACTGTGGGCCGGCAGTGCTGCCATCTGTTGCAAAGCATTACGGATCTAATTACTCTAT CGCTTATCTGCGGGAACTCTCAAAGACTAACAAGCAGGGAACAACAGCTCTTGGCATTG TTGAAGCTGCTAAAAAGTTAGGCTTTGAAACACGCTCTATCAAGGCGGATATGACGCTT TTTGATTATAATGATTTGACCTATCCTTTTATCGTCCATGTGATTAAAGGAAAACGTCT GCAGCATTATTATGTCGTCTATGGCAGCCAGAATAATCAGCTGATTATTGGAGATCCTG ATCCTTCAGTTAAGGTGACTAGGATGAGTAAGGAACGCTTTCAATCAGAGTGGACAGGC CTTGCAATTTTCCTAGCTCCTCAGCCTAACTATAAGCCTCATAAAGGTGAAAAAATGG TTTGTCTAATTTCTTCCCGTTGATCTTTAAGCAGAAAGCTTTGATGACTTATATTATCA TAGCTAGCTTGATTGTGACGCTCATTGATATTGTCGGATCATACTATCTCCAAGGAATA TTGGACGAGTACATTCCTGATCAGCTGATTTCAACTTTAGGAATGATTACGATTGGTCT GATAATAACCTATATTATCCAGCAGGTCATGGCTTTTGCAAAAGAATACCTCTTGGCCG TACTCAGTTTGCGTTTAGTCATTGATGTTATCCTGTCTTATATCAAACATATTTTTACG CTTCCTATGTCTTTCTTTGCGACAAGGCGAACAGGAGAAATCACGTCTCGTTTTACAGA TGCCAATCAGATTATTGATGCTGTAGCGTCAACCATCTTTTCAATCTTTTTAGATATGA CTATGGTAATTTTGGTTGGTGGGGTTTTGTTGGCGCAAAACAATAACCTTTTCTTA ACCTTGCTCTCCATTCCGATTTATGCCATCATTATTTTTGCTTTCTTGAAACCCTTTGA GAAAATGAATCACGAAGTGATGGAAAGCAATGCTGTGGTAAGTTCTTCTATCATTGAAG ATATCAATGGGATGGAAACCATTAAATCACTCACAAGTGAGTCCGCTCGTTATCAAAAC ATTGATAGTGAATTTGTTGATTATTTGGAGAAAAACTTTAAGCTACACAAGTATAGTGC CATTCAAACCGCATTAAAAAGCGGTGCTAAGCTTATCCTCAATGTTGTCATTCTCTGGT ATGGCTCTCGTCTAGTTATGGATAATAAAATCTCAGTTGGTCAGCTTATCACCTTTAAT GCTTTGCTGTCTTATTTCTCAAATCCAATTGAAAATATTATCAATCTGCAATCCAAACT GCAGTCAGCTCGCGTTGCCAATACACGTCTTAATGAGGTCTATCTTGTCGAATCTGAAT TTGAAAAAGACGGCGATTTATCAGAAAATAGCTTTTTAGATGGTGATATTTCGTTTGAA AATCTTTCTTATAAATATGGATTTGGGCGAGATACCTTATCAGATATTAATTTATCAAT CAAAAAAGGCTCCAAGGTCAGTCTAGTTGGAGCCAGTGGTTCTGGTAAAACAACTTTGG CTAAACTGATTGTCAATTTCTACGAGCCTAACAAGGGGATTGTTCGAATCAATGGCAAT GATTTAAAAGTTATTGATAAGACAGCTTTGCGGCGGCATATTAGCTATTTGCCGCAACA GGCCTATGTTTTTAGTGGCTCTATTATGGATAATCTCGTTTTAGGAGCTAAAGAAGGAA CGAGTCAGGAAGACATTATTCGTGCTTGTGAAATTGCTGAAATCCGCTCGGACATTGAA CAAATGCCTCAGGGCTATCAGACAGAGTTATCAGATGGTGCCGGTATTTCTGGCGGTCA AAAACAGCGGATTGCTTTAGCTAGGGCCTTATTAACACAGGCACCGGTTTTGATTCTGG ATGAAGCCACCAGCAGTCTTGATATTTTGACAGAAAAGAAAATTATCAGCAATCTCTTA CAGATGACGGAGAAAACAATAATTTTTGTTGCCCACCGCTTAAGCATTTCACAGCGTAC TGACGAAGTCATTGTCATGGATCAGGGAAAAATTGTTGAACAAGGCACTCATAAGGAAC TTTTAGCTAAGCAAGGTTTCTATTATAACCTGTTTAAT

(A) > S. mutans ComA protein

Figure II (cont'd)

Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al

Atty Dkt No.: 1889-00401

Contact: C.M. Mintz (713)238-8000

Sheet 18 of 19

MKQVIYVVLIVIAVNILLEIIKRVTKRGGTVSSSNPLPDGQSKLFWRRHYKLVPQIDTR
DCGPAVLASVAKHYGSNYSIAYLRELSKTNKQGTTALGIVEAAKKLGFETRSIKADMTL
FDYNDLTYPFIVHVIKGKRLQHYYVVYGSQNNQLIIGDPDPSVKVTRMSKERFQSEWTG
LAIFLAPQPNYKPHKGEKNGLSNFFPLIFKQKALMTYIIIASLIVTLIDIVGSYYLQGI
LDEYIPDQLISTLGMITIGLIITYIIQQVMAFAKEYLLAVLSLRLVIDVILSYIKHIFT
LPMSFFATRRTGEITSRFTDANQIIDAVASTIFSIFLDMTMVILVGGVLLAQNNNLFFL
TLLSIPIYAIIIFAFLKPFEKMNHEVMESNAVVSSSIIEDINGMETIKSLTSESARYQN
IDSEFVDYLEKNFKLHKYSAIQTALKSGAKLILNVVILWYGSRLVMDNKISVGQLITFN
ALLSYFSNPIENIINLQSKLQSARVANTRLNEVYLVESEFEKDGDLSENSFLDGDISFE
NLSYKYGFGRDTLSDINLSIKKGSKVSLVGASGSGKTTLAKLIVNFYEPNKGIVRINGN
DLKVIDKTALRRHISYLPQQAYVFSGSIMDNLVLGAKEGTSQEDIIRACEIAEIRSDIE
QMPQGYQTELSDGAGISGGQKQRIALARALLTQAPVLILDEATSSLDILTEKKIISNLL
QMTEKTIIFVAHRLSISQRTDEVIVMDQGKIVEQGTHKELLAKQGFYYNLFN

## S. mutans comB gene-

ATGGATCCTAAATTTTTACAAAGTGCAGAATTTTATAGGAGACGCTATCATAATTTTGC GACACTATTAATTGTTCCTTTGGTCTGCTTGATTATCTTCTTGGTCATATTCCTTTGTT TTGCTAAAAAAGAAATTACAGTGATTTCTACTGGTGAAGTTGCACCAACAAAGGTTGTA GATGTTATCCAATCTTACAGTGACAGTTCAATCATTAAAAATAATTTAGATAATAATGC AGCTGTTGAGAAGGGAGACGTTTTAATTGAATATTCAGAAAATGCCAGTCCAAACCGTC AAACACCAAAAGAGCAAGAAAAAGAAGAGTCTAAGAGCAAGAAAGCTTCCAAAGATAA GAAAAAGAAATCGAAAGACAAGGAAAGCAGCTCTGACGATGAAAATGAGACAAAAAAGG TTTCGATTTTTGCTTCAGAAGATGGTATTATTCATACCAATCCCAAATATGATGGTGCC AATATTATTCCGAAGCAAACCGAGATTGCTCAAATCTATCCTGATATTCAAAAAACAAG AAAAGTGTTAATCACCTATTATGCTTCTTCTGATGATGTTGTTTCTATGAAAAAGGGGCC AAACCGCTCGTCTTTCCTTGGAAAAAAGGGAAATGACAAGGTTGTTATTGAAGGAAAA ATTAACAATGTCGCTTCATCAGCAACTACTACTAAAAAAGGAAATCTCTTTAAGGTTAC TGCCAAAGTAAAGGTTTCTAAGAAAAATAGCAAACTCATCAAGTATGGTATGACAGGCA AGACAGTCACTGTCATTGATAAAAAGACTTATTTTGATTATTTCAAAGATAAATTACTG CATAAAATGGATAAT

# S. mutans ComB protein

MDPKFLQSAEFYRRRYHNFATLLIVPLVCLIIFLVIFLCFAKKEITVISTGEVAPTKVV DVIQSYSDSSIIKNNLDNNAAVEKGDVLIEYSENASPNRQTEQKNIIKERQKREEKEKK KHQKSKKKKKSKSKKASKDKKKSKDKESSSDDENETKKVSIFASEDGIIHTNPKYDGA NIIPKQTEIAQIYPDIQKTRKVLITYYASSDDVVSMKKGQTARLSLEKKGNDKVVIEGK INNVASSATTTKKGNLFKVTAKVKVSKKNSKLIKYGMTGKTVTVIDKKTYFDYFKDKLL HKMDN



Express Mail No. EL818623952US Inventor(s): Cvitkovitch et al Atty Dkt No.: 1889-00401 Contact: C.M. Mintz (713)238-8000

Sheet 19 of 19

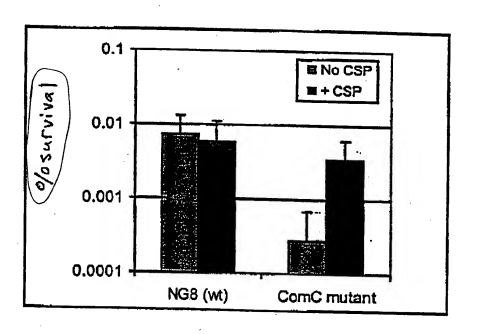


Figure 12